## **PCT**

## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7: WO 00/67558 (11) International Publication Number: A01H 5/00, C07H 21/04, C12N 5/14, **A1** (43) International Publication Date: 16 November 2000 (16.11.00) 15/29, 15/52, 15/82 PCT/US00/12450 (81) Designated States: AE, AG, AL, AM, AT, AU, AZ, BA, BB, (21) International Application Number: BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, (22) International Filing Date: 5 May 2000 (05.05.00) IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, (30) Priority Data: UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, 6 May 1999 (06.05.99) US 60/132,919 GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, (71)(72) Applicant and Inventor: TIMKO, Michael [US/US]; 1610 Old Ballard Road, Charlottesville, VA 22901 (US). CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG). (74) Agent: HANSEN, Christine, M.; Connolly Bove Lodge & Hutz LLP, 1210 Market Street, P.O. Box 2207, Wilmington, DE **Published** 19899 (US). With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: REGULATION OF GENE EXPRESSION IN TOBACCO FOR MANIPULATION OF PLANT GROWTH AND SECONDARY METABOLISM

#### (57) Abstract

This invention relates to enzymes involved in alkaloid, and specifically nicotine, formation in tobacco plants. The invention is based, at least in part, on the nucleotide sequences encoding four variants of putrescine N-methyltransferase (PMT1, PMT2, PMT3, and PMT4), two variants of arginine decarboxylase (ADC1 and ADC2), ornithine decarboxylase (ODC), S-adenosylmethionine synthetase (SAMS), a fragment of NADH dehydrogenase, and a fragment of phosphoribosylanthranilate isomerase. The invention also relates to proteins expressed by these nucleotides, promoter regions of these nucleotides, use of these promoter regions to culture transgenic plant cells and to produce transgenic plants, sense and antisense nucleotides complementary to all or portions of these nucleotide sequences, use of sense and antisense nucleotides to regulate gene expression, and assays using proteins involved in alkaloid formation in tobacco plants.

## FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
$\mathbf{AM}$	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	ТJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JР	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	$\mathbf{z}\mathbf{w}$	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

1

# REGULATION OF GENE EXPRESSION IN TOBACCO FOR MANIPULATION OF PLANT GROWTH AND SECONDARY METABOLISM

## CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a continuation-in-part of US Patent Application Ser. No. 60/132,919, filed May 6, 1999, now abandoned, which is hereby incorporated by reference in its entirety herein.

#### FIELD OF THE INVENTION

This invention relates to enzymes involved in alkaloid, and specifically nicotine, formation in tobacco plants. The invention is based, at least in part, on the nucleotide sequences encoding four variants of putrescine N-methyltransferase (PMT1, PMT2, PMT3, and PMT4), two variants of arginine decarboxylase (ADC 1 and ADC2), ornithine decarboxylase (ODC), S-adenosylmethionine synthetase (SAMS), a fragment of NADH dehydrogenase, and a fragment of phosphoribosylanthranilate isomerase. The invention also relates to proteins expressed by these nucleotides, promoter regions of these nucleotides, use of these promoter regions to culture transgenic plant cells and to produce transgenic plants, sense and antisense nucleotides complementary to all or portions of these nucleotide sequences, use of sense and antisense nucleotides to regulate gene expression, and assays using proteins involved in alkaloid formation in tobacco plants.

20

25

30

5

10

15

## **BACKGROUND OF THE INVENTION**

#### I. Alkaloid Formation

Alkaloids are one of the most diverse groups of secondary compounds found in plants and they are the product of a complex biosynthesis pathway (Hashimoto and Yamada, 1994; Chou and Kutchan, 1998; Waterman, 1998). Why plants accumulate these compounds and in so many different forms is not known. Moreover, for many alkaloids, the exact site of synthesis and the factors that control their intercellular distribution and accumulation remain to be determined (Hashimoto and Yamada, 1994; Kutchan, 1995; Chou and Kutchan, 1998).

Nicotine is the most abundant alkaloid present in cultivated tobacco. Nicotine is formed primarily in the roots of the tobacco plant and subsequently is transported to the leaves, where it is stored (Tso, Physiology and Biochemistry of Tobacco Plants, pp. 233-34, Dowden, Hutchinson & Ross, Stroudsburg, Pa. (1972)).

The synthesis and accumulation of nicotine and other tobacco alkaloids are known to be controlled by various developmental, environmental, and chemical cues. Changes in phytohormone

5

10

15

20

25

30

2

PCT/US00/12450

(e.g., auxin, cytokinin) levels and/or ratios as a consequence of developmental age (Hashimoto and Yamada, 1994; Kutchan, 1995) or by direct manipulation of plant cell culture conditions have been shown to affect the synthesis and accumulation of nicotine and various tobacco alkaloids (Hashimoto and Yamada, 1994; Hibi et al., 1994; Eilbert, 1998). Various abiotic factors (wounding, drought stress, pH imbalance, etc.) [Hashimoto and Yamada, 1994; Kutchan, 1998; Waterman, 1998) 1, 2, 4], as well as biotic factors, such as herbivory, insect feeding, and attack by various microbial and fungal pathogens, are known elicit increased production of nicotine and other alkaloids in the leaves of wild and cultivated tobacco species (Baldwin, 1989; Saito and Murakoishi, 1998; Baldwin and Prestin, 1999). In addition, the commercial practice of topping (i.e., removal of flowering head and young leaves at the upper portions of the plant), results in increases in nicotine and the amount and complexity total alkaloids present in the leaves of Nicotiana tabacum (Hashimoto and Yamada, 1994; Hibi et al., 1994). The factors controlling the topping-induced increase in alkaloid biosynthesis are not known, but likely involve a complex physiological response in the plant as a result of altered phytohormones and wound induced signaling (Akehurst, 1981; Hibi et al., 1994; Kutchan, 1998). In this regard, considerable evidence now exists indicating that a jasmonic acid (JA)- mediated signal transduction pathway may play a role in regulation of gene expression contributing to this increase in alkaloid biosynthesis (Baldwin et al., 1994, 1996, 1997; Ohnmeiss et al., 1997; Imanishi et al., 1998a, 1998b).

The nicotine molecule is comprised of two heterocyclic rings, a pyridine moiety and a pyrrolidine moiety, each of which is derived from a separate biochemical pathway. The pyridine moiety of nicotine is derived from nicotinic acid. The pyrrolidine moiety of nicotine is provided through a pathway leading from putrescine to N-methylputrescine and then to N-methylpyrroline. (Goodwin and Mercer, Introduction to Plant Biochemistry, pp. 488-91, Pergamon Press, New York, (1983)).

Putrescine is formed in plants by one of two pathways (Chattopadhyay and Ghosh, 1998). It can be synthesized directly from ornithine, in a reaction catalyzed by the enzyme ornithine decarboxylase (ODC, EC 4.1.1.17), or formed indirectly from arginine in a reaction sequence initiated by arginine decarboxylase (ADC, EC 4.1.1.19). Putrescine formed by the ADC and/or ODC pathway serves as precursor in the synthesis of the higher polyamines, spermine and spermidine, catalyzed by the enzymes spermine synthase and spermidine synthase, respectively, or it is converted to N-methylputrescine by the action of putrescine N-methyltransferase (PMT), the first committed step in nicotine biosynthesis (Hashimoto and Yamada, 1994; Kutchan, 1995; Chattopadhyay and Ghosh, 1998). N-methyl putrescine is oxidized by a diamine oxidase and cyclized to form the 1-methyl-Δ¹-pyrrolium cation, which is condensed with nicotinic acid or its derivative to form nicotine

3

PCT/US00/12450

(Hashimoto and Yamada, 1994).

Putrescene is a precursor for N-methylputrescine, which then forms N-methylpyrroline. Conversion of putrescine to N-methylputrescine is catalyzed by the enzyme putrescine N-methyltransferase ("PMT"), with S-adenosylmethionine serving as the methyl group donor. PMT appears to be the rate-limiting enzyme in the pathway supplying N-methylpyrroline for nicotine synthesis in tobacco (Feth et al., "Regulation in Tobacco Callus of Enzyme Activities of the Nicotine Pathway", Planta, 168, pp. 402-07 (1986); Wagner et al., "The Regulation of Enzyme Activities of the Nicotine Pathway in Tobacco", Physiol. Plant., 68, pp. 667-72 (1986)).

## II. TRANSGENIC PLANTS

5

10

15

20

25

30

The methods of nicotine formation in tobacco and the genes involved have been studied both to better understand differential gene expression during tobacco growth and development, and also to discover tools useful for creating transgenic plants. For example, the regulatory sequences that modify protein expression in tobacco may be useful in creating transgenic tobacco or other transgenic plants.

It has already been demonstrated that tissues of many plant species may be transformed by exogenous, typically chimeric, genes which are effective to stably transform cells of the tissues. For several species, tissues transformed in this fashion may be regenerated to give rise to whole transquenic or genetically engineered plants. The engineered traits introduced into the transgenic plants by these techniques have proven to be stable and have also proven to be transmissible through normal Mendellian inheritance to the progeny of the regenerated plants. One such desirable trait is the production in the plant cells of desired gene products in vivo in the cells of the transquenic plants. For a chimeric gene to be effective, the foreign DNA sequence containing a coding region should be flanked by appropriate promotion and control regions. Commonly used plant cell transcription promoters include the nopaline synthase promoter from the T-DNA of A. tumefaciens and the 35S promoter from the cauliflower mosaic virus.

In order for the newly inserted chimeric gene to express the protein for which it codes in the plant cell, the proper regulatory signals must be present and in the proper location with respect to the gene. These regulatory signals include a promoter region, a 5' non-translated leader sequence and a 3' polyadenylation sequence. A promoter is a DNA sequence that directs the cellular machinery of a plant to produce RNA from the contiguous structural coding sequence downstream (3') to the promoter. The promoter region influences the rate at which the RNA product of the gene and resultant protein product of the gene is made. The 3' polyadenylation signal is a non-translated region that functions in

5

10

15

20

25

30

4

PCT/US00/12450

the plant cells to cause the addition of polyadenylate nucleotides to the 3' end of the RNA to enable the mRNA to be transported to the cytoplasm and to stabilize the mRNA for subsequent translation of the RNA to produce protein.

Other plant cell transformation techniques are directed toward the direct insertion of DNA into the cytoplasm of plant cells from which it is taken up, by an uncharacterized mechanism, into the genome of the plant. One such technique is electroporation, in which electric shock causes disruption of the cellular membranes of individual plant cells. Plant protoplasts in aqueous solution when subject to electroporation will uptake DNA from the surrounding medium. Another technique involves the physical acceleration of DNA, coated onto small inert particles, either into reqenerable plant tissues or into plant germline cells.

The availability of cloned nucleic acid sequences encoding an enzyme involved in alkaloid synthesis allows for the potential manipulation of alkaloid contents. Furthermore, the availability of promoters useful for expressing genes in plants allows for the creation of chimeric molecules and transgenic plants, which in turn result in possible native plant production of desirable proteins.

Previously reported work discloses cloning nucleotides encoding proteins involved in the biosynthesis of nicotine, and isolating such proteins. Approximately twenty or more cDNAs and/or genomic DNA fragments encoding different enzymes involved with alkaloid formation have been isolated (Chattopadhyay and Ghosh, 1998). For example, successful cloning of partial or full-length cDNA encoding ODC activity from tobacco was disclosed by (Malik et al., J. Plant Biochem. &Biotech. 5:109-112 (1996)). Also, a relatively crude preparation of PMT (30-fold purification) has been subjected to limited characterization (Mizusaki et al., "Phytochemical Studies on Tobacco Alkaloids XIV. The Occurrence and Properties of Putrescine N-Methyltransferase in Tobacco Plants", Plant Cell Physiol., 12, pp. 633-40 (1971)). A process for purifying PMT is disclosed in US Patent No. 5,369,023, "Method of purifying putrescine n-methyltransferase from tobacco plant extract with an anion exchange medium", hereby incorporated by reference in its entirety herein. Several laboratories have reported the cloning of partial or full-length cDNAs encoding ADC (Bell and Malmberg, 1990; Rostogi et al., 1993; Perez-Amador et al., 1995; Nam et al., 1997; Watson and Malmberg, 1996). Comparisons of the amino acid sequences of ADC from various plants revealed a high degree of conservation among the various proteins, as well as homology to ODC (Malmberg et al., 1998).

It is an object of the present invention to characterize the nucleotide and amino acid sequences of enzymes involved in the biosynthesis of nicotine in tobacco. It is also an object of the present invention to provide plant promoter regions that are capable of conferring high levels of transcription in rapidly dividing cells of transformed plants when coupled with a heterologous coding

5

10

15

20

25

30

sequence in a chimeric gene. Further, the invention is directed to chimeric genes incorporating such promoter regions, stable transfection of plants with these chimeric genes, and the plants and cells that are transfected, as well as seeds of such transfected plants. It is a further object to characterize sense and antisense nucleotides capable of regulating expression of genes encoding for enzymes involved in the biosynthesis of alkaloids.

#### **SUMMARY OF THE INVENTION**

Proteins involved in the biosynthesis of nicotine in tobacco *N. tabacum* are the subject of this invention. More specifically, the invention concerns four variants of putrescine N-methyltransferase (PMT1, PMT2, PMT3, and PMT4), two variants of arginine decarboxylase (ADC 1 and ADC2), ornithine decarboxylase (ODC), S-adenosylmethionine synthetase (SAMS), NADH dehydrogenase, and phosphoribosylanthranilate isomerase.

## **BRIEF DESCRIPTION OF THE FIGURES**

Figure 1. Genomic DNA gel blot analysis of the PMT gene family in N. tabacum cv. Xanthi. Total genomic DNA (30 μg) was digested with KpnI, EcoRI, or EcoRI and KpnI, separated by agarose gel electrophoresis, and transferred to nylon membranes. The membrane was hybridized with a <sup>32</sup>P-labeled antisense strand probe covering the complete coding region of the NtPMT1a cDNA. Identity of the hybridizing bands as determined by comparison to phage DNA digests is indicated. Molecular weights are given in kb. Note that KpnI shifts only the NtPMT1b band in the gel blot because this restriction site is present ony in Exon 1 of NtPMT1b and not NtPMT1a.

Figure 2. Amino acid sequence alignment of N. tabacum PMTs. Shown is a PILEUP alignment of the predicted amino acid sequences of the various tobacco PMTs. Amino acid residues that differing among the PMTs are shaded. NtPMT1a, NtPMT2, NtPMT3, and NtPMT4 refer to the deduced amino acid sequences of the PMTs encoded by the NtPMT1a, NtPMT2, NtPMT3, and NtPMT4 genes, respectively, isolated from N. tabacum ev. Xanthi genomic DNA; eNtPMT1a is the predicted amino acid sequence of the A411 cDNA (Accession No. D28506) isolated from N. tabacum ev. Burley 21 by Hibi et al. (1994). The location of the exon-intron boundaries are indicated by the dark vertical line. The nucleotide sequences for NtPMT1a, NtPMT2, NtPMT3, and NtPMT4 appear in GenBank under the accession numbers AF126810, AF126809, AF126811, and AF126812, respectively

Figure 3. Polyacrylamide gel electrophoresis analysis of PCR amplified genomic DNA fragments

encoding Exon 1 of PMT from various species of *Nicotiana*. PCR amplification was carried out as described in the Materials and Methods using Exon 1-specific primers 1 and 2 and total genomic DNA isolated from *N. tabacum*, *N. otophora*, and *N. tomentosiformis*. The amplification products were separated by electrophoresis on 6.5% polyacrylamide gels, the gels fixed, and subject to autoradiography. The amplification products isolated from *N. tabacum* cv. Burley 21 and *N. tabacum* cv. Xanthi were identical and only the amplication products from the reactions with *N. tabacum* cv. Burley 21 DNA are shown. Standards were generated in identical reaction conditions primed with plasmid DNA encoding the various *PMT* genes isolated in this study.

10

15

20

- Figure 4. Nucleotide sequence alignment of the 5'-flanking regions of the N. tabacum PMT genes. Shown is a PILEUP alignment of the nucleotide sequences upstream of the initiating methionine (MET) codon of the four PMT genes isolated from N. tabacum cv. Xanthi. The proposed start site for transcription of the NtPMT1a gene is indicated by the +1 above the sequences. The TATA-box and CCAAT-box motifs are boxed. Potential transcriptional regulatory elements identified by MOTIF search programs are also boxed and indicated by the following abbreviations:. PAL: palindromic sequences; G-Box: G-Box homologous sequences; MRE: metal-responsive element homolog. Nucleotides identical in three or more sequences are shaded. The polyguanine-rich region is underlined. Numbering is indicated to the right and is relative to the proposed start site of each gene.
- Figure 5. RNA gel blot analysis of *PMT* transcript levels in various tissues. Total RNA was isolated from various tissues of mature *N. tabacum* cv. Burley 21 and analyzed by gel blot analysis using a <sup>32</sup>P-labeled *NtPMT1a* cDNA coding region (Exons 2 to 8) probe capable of detecting all *PMT* transcripts.
- 25 A. PMT transcript levels in various tobacco plant tissues and/or organs.
  - B. Induction of *PMT* expression in tobacco roots following topping. Abbreviations: HP, wild-type (*Nic1Nic2*) Burley 21; LP, low alkaloid (*nic1nic2*) mutant. The  $\beta$ -subunit of mitochondrial ATPase ( $\beta$ -ATPase) served as a control.
- 30 Figure 6. Semi-quantitative RT-PCR analysis of PMT gene expression in roots of tobacco plant before and after topping.
  - A. Shown is relative abundance of the individual *PMT* gene transcripts before and after topping. RT-PCR was carried out as described in the Material and methods using Exon 1 specific primers.

    Messenger RNA was amplified from total RNA isolated from the roots of wild-type (HP,

Nic1Nic2) Burley 21 and low alkaloid (LP, nic1nic2) Burley 21 tobacco plants. Far right lane represents size standards for the genes isolated by PCR amplification from plasmid DNA. The β-subunit of mitochondrial ATPase (β-ATPase) served as a control.

- B. Bar graphs showing relative expression of the individual PMT genes following topping in both HP and LP tobacco roots. Abbreviations: HP, wild-type (*Nic1Nic2*) Burley 21; LP, low alkaloid (*nic1nic2*) mutant.
- Figure 7. The nucleotide and predicted amino acid sequences of the transcribed portions of the N.
   tabacum cv Xanthi NtADC1 and NtADC2 genes. Shown are the complete nucleotide and predicted amino acid sequence of the N. tabacum cv Xanthi NtADC1 gene and where it differs from the NtADC2 gene sequence. The dots indicate nucleotide sequence identity and the stars indicate amino acid sequence identity. The proposed polyadenylation signal is underlined. The sequences terminate at the point of polyadenylation found in the full length cDNA (Wang, 1999; AF127239). The
   complete nucleotide sequences for the N. tabacum cv Xanthi NtADC1 (AF127240) and NtADC2 (AF127241) including the 5' and 3' flanking sequences appear in Genbank.
- Fig. 8. Comparison of the predicted amino acid sequences of arginine decarboxylases (ADCs) from various species. Shown is a PILEUP alignment of the predicted amino acid sequence of the N.
  tabacum cv Xanthi NtADC1 gene (AF127240) aligned to the predicted ADC protein sequences from N. sylvestris (AB12873), Arabidopsis thaliana (AF009647), Avena sativa (oat) (X56802), Lycopersicon esculentum (tomato) (L16582) and Escherichia coli (M31770). Amino acid residues conserved among the various ADC are shaded.
- Fig. 9. Gel blot analysis of *ADC* transcript levels in the roots of wild-type and low alkaloid mutant Burley 21 tobacco before and after topping. Total RNA was isolated from the roots of mature wild-type and low alkaloid mutant *N. tabacum* cv. Burley 21 and analyzed by gel blot analysis using [α-<sup>32</sup>P]-dCTP labeled probes recognizing the coding region of ADC or the β-subunit of tobacco mitochondrial ATP synthase (Boutry and Chua, 1985). Quantitation was carried out by phosphorimaging using a Molecular Dynamics PhosphorImager. Values were normalized relative to the intensities of the *atp2* control band in each lane. The experiment was conducted twice with different total RNA samples.

5

10

15

30

8

PCT/US00/12450

- Fig. 10. Differential expression of NtADC-1 and NtADC-2 in various tissues of tobacco. Expression of the NtADC-1 and NtADC-2 genes was analyzed using semi-quantitative RT-PCR and gene specific primers capable of discriminating between transcripts arising from the two genes. Panel A shows control reactions demonstrating primer specificity in the PCR reactions using plasmids containing the NtADC-1 and NtADC-2 coding sequences. The numbers above the lane refer to the specific primer combinations as described in the Materia and methods. Panel B shows the results of RT-PCR reactions using first strand cDNA synthesized from total RNA extracted from either root, leaf, or flowers. As a internal control, primers specific for the *atp2* gene transcript were include in the amplification reactions. All reactions were carried out within the linear range of template amplification as determined by varying template amount, amplification time, and temperature as described in Riechers and Timko (1999).
- Fig. 11. Genomic DNA gel blot analysis of the ODC gene family in *N. tabacum*. Total genomic DNA (30  $\mu$ g) was digested with *Eco*RI or *Hind*III, fractionated by agarose gel electrophoresis, transferred to nylon membranes and hybridized with an  $\alpha$ -<sup>32</sup>P-dCTP labeled probe encoding full-length ODC cDNA as described in the Materials. The mobility of molecular weights standards are given to the right of the figure in kilobases (kb).
- Fig 12. Comparison of the nucleotide and predicted amino acid sequences of the NtODC-1 and NtODC-2 genes. Shown are the nucleotide and predicted amino acid sequences of the NtODC-1 (AF233850) and NtODC-2 (AF233849) genes. In the figure, the complete amino acid sequence of the pODC2 is given and the pODC1 sequence is given only where it differs. The start site of transcription is designated as +1 and the poly(A) addition site is indicated by the arrow. Within the relevant regions of homology, nucleotide differences between the NtODC-1 and NtODC-2 genes are in bold lettering. The proposed TATA-box, and polyadenylation signal are shaded.
  - Fig. 13. Protein sequences alignment of ornithine decarboxylases (ODCs) from various species. Shown is a PILEUP alignment of the predicted amino acid sequences of the *N. tabacum* cv. Xanthi pODC2 protein (AF233849) with the ODCs from *N. tabacum* cv. SC58 (Y10472) and cv. BY-2 (ABO31066), *Lycopersicon esculentum* (tomato) (AF030292), *Datura stramonium* (jimsonweed) (X87847), *Saccharomyces cerevisiae* (NP\_012737), and humans (*Homo sapiens*; AAA59966). Amino acid residues conserved among the various ODCs are shaded.

9

PCT/US00/12450

**Fig. 14.** Gel blot analysis of *ODC* transcript levels in various tissues of mature tobacco plants and in the roots before and after topping. Total RNA was isolated from various tissues of mature N. *tabacum* cv. Burley 21 and analyzed by gel blot analysis using an α- $^{32}$ P-dCTP labeled coding region probes for ODC. (A) Transcript levels in various organs of wild-type tobacco: R, root: S, stem; L, leaf; SE, sepal; PE, petal; O, ovary; S, stamen; and AN, anther. (B) Transcript levels in roots of Burley 21 tobacco plants before and after topping. RNA gel blot analysis of the tissues-specific distribution and post-topping expression of transcripts encoding ODC in tobacco. As a control, the blots were also probed with radioactively labeled probes encoding the alkaloid biosynthesis enzyme putrescine N-methyltransferase (PMT) and a root specific β-glucosidase (TBG-1).

10

15

20

25

30

5

## **DETAILED DESCRIPTION OF THE INVENTION**

Nucleic acid sequences have been isolated from tobacco that encode important enzymes in nicotine and total alkaloid formation, including PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, and SAMS. Also identified are cDNA fragments encoding partial segments of NADH dehydrogenase and phosphoribosilanthronilate isomerase. Also identified are promoter regions for the nucleotides encoding PMT1, PMT2, PMT3, PMT4, and ADC2. All of these nucleic acids are isolated from *Nicotiana tabacum* L.

"Promoter" and "promoter region" are terms used interchangeably herein to refer to a DNA sequence that regulates expression of a selected DNA sequence operably linked to the promoter, and which effects expression of the selected DNA sequence in cells. The term also encompasses the 5'untranslated region that may be transcribed into mRNA but is not translated.

"Protein", "polypeptide", and "peptide" are used interchangeably herein when referring to a gene product.

In one aspect, the invention features isolated nucleic acid molecules encoding for PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, and SAMS, a fragment of NADH dehydrogenase and a fragment of phosphoribosilanthronilate isomerase. The disclosed molecules can be non-coding (e.g. probe, antisense or ribozyme molecules) or can code for a functional enzyme. In one embodiment, the nucleic acid molecules can hybridize to the nucleic acid sequences encoding for PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, a fragment of NADH dehydrogenase, or a fragment of phosphoribosilanthronilate isomerase or to the complements of these nucleic acid sequences. In a preferred embodiment, the hybridization is conducted under mildly stringent or stringent conditions.

In further embodiments, the nucleic acid molecule is at least 50%, 60%, 70%, 80% and more preferably at least 90% or 95% homologous in sequence to the nucleic acid sequences encoding for PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, a fragment of NADH dehydrogenase, or

a fragment of phosphoribosilanthronilate isomerase or to the complements of these nucleic acid sequences. In another embodiment, the nucleic acid encodes a polypeptide that is at least 50%, 60%, 70%, 80% and more preferably at least 90% or 95% similar in sequence to the amino acid sequence of PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, the fragment disclosed herein of NADH dehydrogenase, or the fragment of phosphoribosilanthronilate isomerase disclosed herein.

5

10

15

20

25

30

In another embodiment, the invention features isolated polypeptides, preferably substantially pure preparations, encoded for by the nucleic acid sequences of the invention. Particularly preferred are those polypeptides encoded for by the nucleic acid sequences identified by (SEQ. ID. NO. 2), (SEQ. ID. NO. 5), (SEQ. ID. NO. 8), (SEQ. ID. NO. 11), (SEQ. ID. NO. 13), (SEQ. ID. NO. 15), (SEQ. ID. NO. 18), (SEQ. ID. NO. 21), (SEQ. ID. NO. 23), (SEQ. ID. NO. 25) or (SEQ. ID. NO. 26) or comprising a nucleotide sequence encoding the amino acid sequence encoded by (SEQ ID NO. 3), (SEQ. ID. NO. 6), (SEQ ID. NO. 9), (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) or (SEQ. ID. NO. 24). In particularly preferred embodiments, the subject polypeptides can aid in regulating the production of alkaloids, particularly nicotine, in plants. In one embodiment, the polypeptide is identical to or similar to the protein represented by the amino acid sequences of (SEQ ID NO. 3), (SEQ. ID. NO. 6), (SEQ ID. NO. 9), (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) or (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) or (SEQ. ID. NO. 24). In a preferred embodiment, the polypeptide is encoded by a nucleic acid that hybridizes with a nucleic acid represented in.

The polypeptides of the present invention can comprise full length proteins, such as represented by (SEQ ID NO. 3), (SEQ. ID. NO. 6), (SEQ ID. NO. 9), (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) and (SEQ. ID. NO. 24), or can comprise one or more fragments corresponding to one or more particular motifs/domains, or to arbitrary sizes, e.g., at least 5, 10, 25, 50, 100, 150, or 200 amino acids in length.

Another aspect of the invention features chimeric genes comprised of a promoter for the genes for PMT2, PMT1, PMT3, PMT4, or ADC2. Yet another aspect of the invention features chimeric genes or chimeric molecules comprised respectively of the functional gene encoding for or the protein PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, NADH dehydrogenase and/or phosphoribosilanthronilate isomerase.

The invention also concerns isolated and purified promoter regions for tobacco Betaglucosidase and their use in chimeric genes or chimeric molecules.

Another aspect of the invention involves vectors capable of transporting another nucleic acid to which a vector has been linked. Preferably, the vectors comprise the nucleic acid sequences of the invention or their complements.

5

10

15

20

25

30

11

The invention also features transgenic plants and their seeds that include (and preferably express) a heterologous form of PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, NADH dehydrogenase and/or phosphoribosilanthronilate isomerase. The present invention also encompasses transgenic plants that contain in their genome a chimeric gene construction incorporating the nucleic acid encoding PMT1, PMT2, PMT3, PMT4, ADC1, ADC2, ODC, SAMS, NADH dehydrogenase and/or phosphoribosilanthronilate isomerase. Such transgenic plants and their seeds may be useful to natively produce enhanced quantities of desirable exogenous proteins, such as compounds useful for pharmaceutical purposes, or proteins that provide herbicide resistance.

PCT/US00/12450

Another feature of the invention is the use as probes of the DNA sequences disclosed herein or oligonucleotide fragments thereof. Probes may be useful to obtain additional gene family members or locate homologous genes in tobacco or other plant species. Copies of related genes can be obtained from existing genomic libraries or the genomic libraries can be constructed. In one embodiment, an isolated DNA sequence comprising about a fifteen to about a twenty-five base pair oligonucleotide sequence identical to any consecutive about fifteen to about twenty-five base pair sequence found in the sequences of the invention is used as a probe.

Another feature is use of the polypeptides of the invention in an assay, such as an assay to identify modulators of enzyme activity in plants.

Other features and advantages of the invention will be apparent to those of skill in the art.

The nucleotide and amino acid sequences of the invention are disclosed herein in the Sequence Listing, text, and the figures. Specific sequences of the invention are provided in the attached Sequence Listing and can be understood to represent promoters, nucleic acids, and proteins respectively relating to the following proteins: PMT2 (SEQ. ID. NOS. 1, 2, and 3); PMT1 (SEQ. ID. NOS. 4, 5, and 6); PMT3 (SEQ. ID. NOS. 7, 8, and 9); PMT4 (SEQ. ID. NOS. 10, 11, and 12); SAMS (SEQ. ID. NOS. 13 and 14); ODC (SEQ. ID. NOS. 15 and 16); ADC1 (SEQ. ID. NOS. 17, 18, and 19); ADC2 (SEQ. ID. NOS. 20, 21, and 22); ADC1 mRNA (SEQ. ID. NOS. 23 and 24); NADH dehydrongenase (SEQ. ID. NO. 25); and PAI (SEQ. ID. NO. 26). If only two sequence identifiers are provided for a protein, those sequences represent the nucleic acid and the protein respectively. If three identifiers are provided, the identifiers represent promoter, genomic or cDNA nucleic acid, and peptide sequences, respectively. If only one identifier is provided, it represents a DNA fragment coding for the protein or portions of it.

For other reference, the sequences may be found at the following records in GenBank at the following Accession Numbers, which records are hereby incorporated in their entirety herein: AF126810 (NtPMT1); AF126809 (NtPMT2); AF126811 (NtPMT3); AF126812 (NtPMT4), AF176908 (NtomPMT)(Nicotiana tomentosiformis); AF76909 (NotoPMT)(Nicotiana otophora);

AF127239 (ADC); AF127240 (ADC1); AF127241 (ADC2); AF127242 (ODC); AF233849 (ODC2); AF233850 (ODC1); and AF127243 (SAMS).

12

PCT/US00/12450

The following experimental discussion is presented to better illustrate the invention.

## I. PMT

5

10

15

20

25

30

WO 00/67558

The present invention features the characterization of four members of the nuclear gene family encoding PMT in tobacco N. tabacum. The nucleic acid sequences encoding PMT and the amino acid sequences for the proteins are reported herein and can also be found in the DDBJ, EMBL, and GenBank Nucleotide Sequence Databases under the accession numbers for NtPMT1a, NtPMT2, NtPMT3, and NtPMT4 at AF126810, AF126809, AF126811, and AF126812, respectively. The complete coding region and immediate 5'- and 3'- flanking regions are characterized.

As the discussion below shows, all four PMT genes present in the N. tabacum genome are expressed in the roots of wild-type plants and differentially regulated in tobacco lines expressing either high or low total alkaloid contents.

#### Materials and Methods

#### Plant materials

Seeds of N. sylvestris, N. otophora, and N. tomentosiformis were obtained from the USDA-ARS national tobacco germplasm collection (Oxford, NC). N. tabacum ev. Burley 21 and N. tabacum ev. Xanthi seeds were kindly provided by Glenn Collins, University of Kentucky. Tobacco plants used for DNA isolation were grown in a soil:vermiculite mixture in the greenhouse under natural lighting conditions. Plants used for RNA extraction were grown in Moltan Plus (Moltan Co., Middleton, TN).

Gel blot analysis of genomic DNA

Young leaves were collected from greenhouse grown tobacco (N. tabacum ev. Xanthi) plants and total genomic DNA was isolated from freshly-harvested tissues using a modification of the CTAB extraction method (Dellaporta et al., 1983). Approximately 30 µg of total DNA was digested with EcoRI, KpnI, or EcoRI and KpnI and the digestion products separated by electrophoresis through a 0.75% agarose gel. Restricted and size-fractionated DNA was denatured and transferred to Nytran+ nylon membranes (Schleicher and Schuell, Keene, NH) by capillary blotting in 0.4N NaOH overnight. Membranes were prehybridized in 0.25M Na<sub>2</sub>HPO<sub>4</sub> (pH 7.4), 7% SDS, 1 mM Na<sub>2</sub>EDTA

13

PCT/US00/12450

for at least 2 hr, then hybridized overnight at 65°C in the same buffer with 2-3 x 10<sup>6</sup> cpm/mL of a <sup>32</sup>P-labeled single-stranded probe (antisense DNA strand). The probe was prepared by the method of Bednarczuk *et al.* (1991) using a primer (Table 1, primer 4) designed from the 3' end of the *NtPMT1a* coding region (Exon 8) and the full-length coding region of the *NtPMT1a* cDNA as template. The *NtPMT1a* cDNA was generated by RT-PCR using synthetic oligonucleotide primers based on the N- and C-terminal sequences of the A411 cDNA reported by Hibi *et al.* (1994) and RNA template isolated from *N. tabacum* cv. Burley 21 roots. Membranes were washed at a final stringency of 0.1 x SSC, 0.1% SDS at 65°C. Hybridizing bands were visualized by autoradiography and/or imaged using a Molecular Dynamics PhosphorImager (Model 445 SI, Sunnyvale, CA).

10

15

20

25

30

5

Genomic library construction and phage isolation

A library of *N. tabacum* cv. Xanthi genomic DNA fragments constructed in EMBL3 was purchased from Clontech (Palo Alto, CA) and a total of 1.1 x 10<sup>6</sup> recombinant phage were screened by plaque hybridization using random-primed <sup>32</sup>P-labeled *NtPMT1a* cDNA as probe (Sambrook *et al.*, 1989). Prehybridization, hybridization, and washing conditions were as described above. Positive hybridizing phage were plaque purified by subsequent rounds of rescreening and DNA was prepared from 18 independently isolated phage. The phage DNA was characterized by restriction analysis and DNA gel blot analysis and fragments containing the sequences encoding PMT were subcloned into pBluescript KS vectors for further analysis.

Comparison of the hybridizing fragments present in the 18 recombinant phage to the hybridization pattern obtained by genomic DNA blot analysis indicated that only three of the *PMT* genes suspected to be present in the *N. tabacum* genome were recovered from the library screen. To obtain sequences encoding NtPMT1a, a subgenomic library was constructed from *N. tabacum* cv. Xanthi DNA. The library consisted of gel-purified 2.5-3.5 kb *Eco*RI fragments ligated into  $\lambda$ \_ZAP II vector arms and packaged using Gigapack III Gold packaging extracts according to the manufacturer's instructions (Stratagene, La Jolla, CA). The primary library was amplified once in *E. coli* XL1-Blue MRF' strain (Stratagene) and screened as described above, except that a random-primed  $^{32}$ P-labeled *NtPMT1a* cDNA Exon 1-specific probe was used (Table 1). Exon 1 had previously been amplified by PCR using primers 1 and 2 (Table 1) and the *NtPMT1a* cDNA as template. The recombinant phage that hybridized with the probe was isolated from the sublibrary by two more rounds of plaque purification, and the pBluescript phagemid containing the approximate 3.1 kb *Eco*RI genomic fragment with the *NtPMT1a* gene was excised from the  $\lambda$ \_ZAP II phage vector using the *in vivo* excision protocol described by Stratagene.

#### DNA sequence analysis

5

10

15

20

25

30

Unless otherwise noted, DNA sequencing was performed with double-stranded plasmid DNA templates using fluorescent dye terminator technology (dRhodamine Terminator Cycle Sequencing Ready Reaction kit) on an ABI 310 DNA sequencer (Perkin-Elmer Applied Biosystems). For analysis of PCR products, following electrophoretic separation of amplification reaction products, the bands of interest were excised from the polyacrylamide gels, the DNA extracted using the Quiagen Gel Extraction Kit, and the recovered DNA used as sequencing template. Sequencing was performed using AmpliTaq DNA polymerase and fluorescent dye terminator technology (as described above) and primers 1 and 2 (Table 1) specific for Exon 1. Nucleotide and amino acid sequences were analyzed and aligned using either the Clustal method and Lasergene software (DNAStar Inc., Madison, WI) or the PILEUP and ALSCRIPT (Genetics Computer Group, Madison, WI) sequence analysis package (Version 9.0). Transcription factor binding site homologies were identified in promoter DNA sequences by searching the transcription factor database using the GCG program.

#### RNA gel blot analysis

For RNA analysis, roots and other tissues were harvested from mature wild-type (HP; *Nic1Nic2*) and low alkaloid mutant (LP; *nic1nic2*) Burley 21 tobacco plants. For topping experiments, the stem was cut and the top one-third of the plant was removed just prior to flower opening. Roots were harvested just prior to topping (0 hr control) and at various times after decapitation. The tissue was immediately frozen in liquid nitrogen and stored at -80°C until RNA extraction and isolation.

Total RNA was isolated from vegetative organs and floral structures of HP and LP Burley 21 tobacco using the TRI-reagent (Molecular Research Center Inc., Cincinnati, OH) and quantified spectrophotometrically by measuring *A*260. Total RNA (5 μg) was electrophoresed through 1.2% agarose gels (containing 0.4 M formaldehyde) and transferred to Nytran+ nylon membranes. Following prehybridization the membranes were hybridized with a single-stranded *NtPMT1a* cDNA antisense probe (corresponding to the antisense strand of Exons 2 to 8 of the *NtPMT1a* cDNA coding region) as described above. As a control to quantify and normalize RNA levels in each lane, the blot was hybridized with a 400-bp probe derived from the β-ATPase cDNA using primers 6 and 7 (Table 1) as described below.

Semi-quantitative RT-PCR analysis of individual PMT transcript levels

Total RNA (1 µg) extracted from the roots of HP and LP Burley 21 tobacco plants was reversetranscribed into first-strand cDNA at 42°C using Superscript II reverse transcriptase (Gibco BRL) according to the manufacturer's protocol. Two gene-specific primers were employed in the reactions: primer 5 capable of recognizing Exon 3 of the PMT genes and primer 8 specific for Exon 8 of the nuclear gene encoding the β-subunit of mitochondrial ATPase from N. plumbaginifolia (NpATP2.1) and N. sylvestris (NsATP2.1) (Boutry and Chua, 1985; Lalanne et al., 1998). The β-ATPase transcript served as an internal reference (constitutively-expressed control) to determine loading accuracy and to normalize expression levels (Kinoshita et al., 1992) Following first strand cDNA synthesis, two sets of nested primers (0.4  $\mu$ M each primer) were used to amplify the PMT and  $\beta$ -ATPase transcripts: primers 1 and 2 (Table 1) recognized Exon 1 in all five PMT transcripts and gave products ranging in size from 220 bp to 420 bp and primers 6 and 7 amplified an approximately 400-bp region encompassing a portion of Exons 6 to 8 of the β-ATPase coding region. Amplification was carried out for 25 cycles using the following reaction conditions: denaturation at 95°C for 1 min, primer annealing at 60°C for 35 sec, and extension at 72°C for 1.5 min; a final extension was conducted at 72°C for 6 min. Amplification products were radioactively labeled by spiking the PCR reaction with 10 µCi 32P-dCTP. Aliquots of the PCR reaction were analyzed on a 6.5% non-denaturing polyacrylamide/1X TBE gel and electrophoresed at 600 volts. The reaction

extension was conducted at 72°C for 6 min. Amplification products were radioactively labeled by spiking the PCR reaction with 10 μCi 32P-dCTP. Aliquots of the PCR reaction were analyzed on 6.5% non-denaturing polyacrylamide/1X TBE gel and electrophoresed at 600 volts. The reaction conditions were optimized to provide amplification of both PMT and \$-ATPase transcripts in the linear range of the reaction by varying the levels of first strand cDNA template, annealing temperature, and number of cycles of amplification as described in Kinoshita *et al.* (1992). Molecular weight standards were prepared by PCR amplification using the same primers and protocol described above and plasmid DNA templates containing the PMT encoding genomic fragments, as well as genomic DNA from the various *Nicotiana* species indicated in the text.

Following electrophoresis, the polyacrylamide gels were fixed in 5% MeOH, 7.5% acetic acid for 30 min, dried overnight, and used to expose X-ray film. PMT band intensities were quantified using phosphorimager analysis (Molecular Dynamics) and normalized relative to the intensities of the  $\beta$ -ATPase control band in each lane. The experiment was conducted twice with different total RNA samples, and representative results are presented from one of the two experiments.

#### Results

5

10

15

20

25

30

PMT gene structure and organization in N. tabacum

5

10

15

20

25

30

Gel blot analysis of total genomic DNA isolated from *N. tabacum* cv. Xanthi, hybridized with a radioactively-labeled cDNA (*NtPMT1a*) encoding the complete coding region of putrescine N-methyltransferase (PMT) showed the presence of five major hybridizing bands in *Kpn*I or *Eco*RI digested DNA, consistent with the presence of a small multigene family in the *N. tabacum* genome (Figure 1).

16

PCT/US00/12450

As part of our initial characterization of the gene family encoding PMT in N. tabacum, an EMBL3 genomic library, prepared from N. tabacum cv. Xanthi DNA, was screened using the NtPMT1a (A411 homologous) cDNA as probe. From a total of 18 recombinant phage isolated, three phage were recovered that contained genomic fragments encoding the NtPMT2, NtPMT3 and NtPMT4 genes. The three PMT genes were completely encoded within a unique sized EcoRI fragment within the phage DNA insert which allowed for the correlation of each with a hybridizing restriction fragment on the gel blot of N. tabacum genomic DNA (Figure 1). The complete coding region and immediate 5' and 3' non-coding sequences of the three genes were determined and found to encode full-length PMT proteins (Figure 2). Each PMT gene consisted of 8 exons and 7 introns, consistent with the gene structure reported previously for the PMT genes from N. sylvestris (Hashimoto et al., 1998a). Comparison of the deduced amino acid sequences (Figure 2) revealed that the encoded PMT proteins were extremely similar over their entire length, with the only significant variability in primary sequence localized to the extreme N-terminal region of the protein. This region, completely encoded within Exon 1, contains a variable number of an 11 amino acid repeat with a consensus sequence of NGHQNGTSEHQ. The function of the repeated sequence is unknown, but is apparently inconsequential to enzyme function, since the number of repeats does not influence activity and PMTs characterized from other species do not contain the repeated element (Hashimoto et al., 1998a; Suzuki et al., 1999a).

Multiple rounds of screening of the EMBL3 genomic library failed to yield additional hybridizing phage containing sequences encoding the other two *PMT* genes thought to be present in the *N. tabacum* genome and, therefore, a directed cloning approach was pursued using a subgenomic library constructed from *EcoRI* fragments isolated from *N. tabacum* cv. Xanthi. From this hybridization screening, a phage containing the approximately 3.1 kb EcoRI fragment encoding *NtPMT1a* was recovered. The coding region of the *NtPMT1a* gene was found to be identical to the A411 cDNA (Hibi *et al.*, 1994), with the exception of a single base change in Exon 6 that results in a conservative amino acid substitution. This difference could be the result of minor differences among cultivars used in the two studies (i.e., Xanthi vs. Burley 21). Translation of the open reading frame contained in *NtPMT1a* showed that it encoded a protein containing four N-terminal 11 amino acid repeats, similar to Exon 1 of the *PMT* gene present in *N. tomentosiformis* (Hashimoto *et al.*, 1998a).

PCT/US00/12450

WO 00/67558

Given the observation that NtPMT1a encoded a homolog of the PMT gene present in N. tomentosiformis, the nature and possible evolutionary origin of the remaining PMT gene present in the N. tabacum genome was brought into question. From our expression studies (described in detail below), we had determined that five distinct PMT encoding transcripts were present in the roots of N. tabacum, four of which could be accounted for based upon the length of the Exon I coding region in the four PMT genes isolated and characterized in our studies described above. The fifth transcript was similar in size to that encoded by NtPMT1a and, therefore, was designated NtPMT1b. Since the variability in PMT gene structure is primarily localized within Exon 1, we used a PCR-based strategy to analyze the PMT gene structure and family size in N. otophora, the other possible progenitor of N. tabacum. As shown in Figure 3, five distinct PCR products were detected in the electrophoretic pattern of amplification products generated from N. tabacum genomic DNA using Exon 1 specific primers (Table 1). Consistent with our studies described above and the previous work of Hashimoto et al. (1998a), three PCR products were detected in the electrophoretic pattern of amplification products generated from N. sylvestris genomic DNA, and a single band was recovered from N. tomentosiformis genomic DNA. Amplification of genomic DNA from N. otophora using Exon 1 specific primers also yielded only a single band, whose electrophoretic mobility was most similar to that of the NtPMT1b derived product.

### Analysis of PMT gene intron and flanking sequences

20

25

30

5

10

15

The location of the seven introns within the protein coding region of the five *PMT* genes in *N. tabacum* is identical and appears to be conserved among *PMT* genes from different *Nicotiana* species. There is also little variation in the nucleotide sequences that comprise the Exon-Intron splice junctions in the various *PMT* genes in *N. tabacum* (Table 2). The high degree of nucleotide sequence similarity recognized among *PMT* genes within their coding regions is also present within their introns and immediate 5' and 3' flanking sequences (Table 2 and Figure 4). In general, a greater level of sequence identity is found in the introns of the *NtPMT2*, *NtPMT3*, and *NtPMT4* genes, than in pair-wise comparisons among the introns of the other members of the *N. tabacum PMT* gene family. The observed conservation in the intron sequences of the *NtPMT2*, *NtPMT3*, and *NtPMT4* genes is consistent with their origin from the same progenitor species (*N. sylvestris*). One interesting exception occurs within Intron 6, where the length of the intron and the sequence similarity is more conserved between *NtPMT1a* and *NtPMT4*, than between *NtPMT4* and *NtPMT2* or *NtPMT3*.

Approximately 1 kb of nucleotide sequence was determined 5' to the coding regions of the *NtPMT1a*, *NtPMT2*, *NtPMT3*, and *NtPMT4* genes (Figure 4). By comparison to the 5'-untranslated

region (UTR) present in the A411 cDNA, we set the start site for transcription initiation at approximately 57 nucleotides upstream of the MET start codon in *NtPMT1a* and *NtPMT3*, and either 69 or 60 nucleotides upstream in *NtPMT2* and *NtPMT4*. The major distinguishing feature between the 5'-UTRs in the various genes is the presence or absence of a 17 bp sequence in the gene. An appropriately placed TATA-box can be easily recognized 45 bp 5' to the initiation site in all four genes. Within the first 200-250 bp upstream of the TATA box, a high level of sequence conservation is found to exist among the promoter regions in the four genes. After this point, a clear difference can be observed between the *NtPMT1a* promoter and the remaining three genes, and by 400 bp upstream, little similarity can be found among any of the gene family members.

Analyzing the proximal regions of the various *PMT* promoters with various motif scanning software identified several G-box-like sequences (Foster *et al.*, 1994; Kim *et al.*, 1992; Menkens *et al.*, 1995; Staiger *et al.*, 1989; Williams *et al.*, 1992) at various positions among the *PMT* promoters, and a potential metal response element (MRE) (positions –75 to –66; numbering relative to the *NtPMT1a* promoter sequence) in three of the four *PMTs* (Cizewski-Culotta and Hamer, 1989; Thiele, 1992). An unusual 17 nucleotide stretch of guanine occurs at positions -259 to -243 in the *NtPMT1a* gene promoter followed upstream by a purine-rich region (positions –332 to –263). In the *NtPMT3* promoter a 14 bp palindromic sequence (positions -497 to -484) was detected. *PMT* gene expression has been reported to increase in root tissues following treatment with methyl jasmonate (Imanishi *et al.*, 1998). However, none of the sequence motifs reported to confer methyl jasmonate-responsiveness in other plant genes (Mason *et al.*, 1993; Rouster *et al.*, 1997) were detected in the *PMT* promoters.

Comparison of the available nucleotide sequence information from the 3'-flanking regions of the various *PMT* genes in *N. tabacum* revealed that the 3'-UTRs in the *NtPMT2*, *NtPMT3*, and *NtPMT4* genes of *N. tabacum* share approximately 81-94% identity with each other and are essentially identical to those reported for *N. sylvestris* PMTs by Hashimoto *et al.* (1998a). The major distinguishing feature among the various genes is the presence of two short (20 bp and 4 bp) deletions in the *NtPMT2* gene, which lowers the percent identity. The 3'-UTR of *NtPMT1a* is identical to that reported for the A411 cDNA (Hibi *et al.*, 1994) and 81-94% identical to the other *PMT* genes in the *N. tabacum* genome. Unfortunately, no sequence information is currently available for the 3'-UTR of the *N. otophora* or *N. tomentosiformis PMT* genes.

## Regulation of PMT gene expression

5

10

15

20

25

30

To determine whether the members of the PMT gene family in N. tabacum were differentially

19

expressed, a series of experiments were carried out to define the temporal and spatial distribution of transcripts arising from the five genes. Shown in Figure 5A are the results of gel blot analysis of total RNA extracted from various tissues of mature Burley 21 tobacco plants hybridized with radioactively-labeled probe capable of detecting all five *PMT* transcripts. Consistent with previous studies (Hashimoto *et al.*, 1998b; Hibi *et al.*, 1994), *PMT* expression is localized exclusively to roots. When maturing wild-type (HP) Burley 21 plants are topped (i.e., the floral meristem and upper 1/3 of the stem are removed), a dramatic increase in *PMT* transcript abundance is observed within 2 hr, reaching a maximal level of accumulation by 12-24 hr. Two size transcripts are detected on the gel blots, reflecting the small difference in message size that occurs as a result of the difference in size of Exon 1 among the genes.

5

10

15

20

25

30

In addition to examining *PMT* gene expression in wild-type plants, we also examined expression in a low nicotine-producing (LP) mutant of Burley 21 (Legg and Collins, 1971). The low nicotine Burley 21 line harbors mutations at two independent loci (*nic1* and *nic2*) thought to be global regulators of gene expression involved in alkaloid formation. As shown in Figure 6B, topping of the low nicotine mutant (*nic1nic2*) Burley 21 did not cause an increase in *PMT* transcript abundance as observed in wild type plants. Thus, it appears that *Nic1* and *Nic2* are likely involved in regulation of *PMT* expression in the very least, and may also be involved in the regulation of other genes in the alkaloid biosynthetic pathway. Whether this is a direct effect (e.g., transcriptional activation) or indirect remains to be determined.

In order to determine the extent to which the individual members of the gene family contributed to the general pattern of expression described above, a semi-quantitative RT-PCR strategy (Kinoshita *et al.*, 1992) was used to detect and quantify the levels of the individual *PMT* transcripts in the roots of both wild-type (HP) and low alkaloid (LP) Burley 21 tobacco. This approach takes advantage of the fact that Exon 1 is variable in length among the various *PMT* genes (Figure 2), allowing for their individual detection and quantitation following polyacrylamide gel electrophoresis and autoradiography.

Five RT-PCR products (representing Exon 1 from each of the five genes present in *N. tabacum*) were detected in the electrophoretic profiles of amplification products derived from reactions using either HP or LP Burley 21 root RNA (Figure 6A). All five *PMT* genes present in the *N. tabacum* genome were expressed in the roots of wild-type plants, and topping resulted in a differential accumulation of transcripts derived from each gene. Among the five genes, transcripts derived from the *NtPMT2* and *NtPMT1b* showed the greatest increase in abundance rising approximately 3-fold during the first 24 hr post-topping, whereas levels of the *NtPMT1a* and *NtPMT4* transcripts changed little in response to topping (Figure 6B). In the LP mutant, little or no effect was observed on the

20

PCT/US00/12450

levels of the various *PMT* transcripts following topping, although in some cases (e.g., *NtPMT1a*) a small but likely insignificant depression in transcript abundance was detected. Thus, it appears that all five genes contribute to PMT activity levels within the root.

## 5 II. ADC

The present invention features the characterization of two members of the nuclear gene family encoding ADC in tobacco *N. tabacum* L. As the following discussion shows, *ADC2* is preferentially expressed in roots and accounts for the major portion of *ADC* transcripts present. Furthermore, analysis of *ADC* transcript levels in roots of low and high nicotine producing lines showed that *ADC* expression is under the control of the *Nic1 Nic2* regulatory loci.

#### Materials and methods

#### Plant growth and tissue preparation

20

15

10

Seeds of *N. tabacum* cv. Xanthi, wild-type and low alkaloid *nic1 nic2* mutant *N. tabacum* cv. Burley 21 were obtained from Dr. G. Collins (University of Kentucky, Lexington). Tobacco plants used for DNA isolation were grown in soil:vermiculite mixture in the greenhouse under natural lighting conditions. Plants used for RNA extraction were grown either in Moltan Plus (Moltan Co., Middleton, TN) or hydoponically in a dilute (half-strength) Peters nutrient solution with continuous aeration of the roots under natural lighting conditions in the greenhouse. Topping experiments were conducted by removing the floral meristem, leaves and stem (approximately the upper 1/3 of the plant) from tobacco plants just prior to blooming. Plant tissues were collected from fully matured individuals, frozen in liquid nitrogen, and stored at -80°C until used for RNA preparation (see below).

Screening of genomic libraries and phage characterization

30

25

A genomic library constructed in  $\lambda$  EMBL3 from *N. tabacum* cv. Xanthi leaf DNA (Clonetech, Inc., Palo Alto, CA) was screened by plaque hybridization (Sambrook *et al.*, 1989) using an [ $\alpha$ -  $^{32}$ P]- dCTP-labeled, 2.7 kb *Eco*RI-*Xho*I fragment from plasmid PR24 as probe. PR24 encodes a full length ADC cDNA isolated from the roots of wild-type *N. tabacum* cv. Burley 21 (Wang, 1999). Hybridization was performed at 65 °C for 16 h in a solution containing 0.25 M Na<sub>2</sub>HPO<sub>4</sub> (pH 7.2) and 7% (w/v) SDS. Following hybridization, the membranes were washed twice in 2 x SSC, 0.1%

SDS for 15 min at room temperature, once in 0.2 x SSC, 0.1% SDS for 30 min at 65°C. Hybridizing phage were picked and plaque purified through three subsequent rounds of hybridization screening. Phage DNA was isolated from plaque purified phage using a Qiagen Phage Midi Preparation Kit (Qiagen, Germany) and insert DNA characterized by restriction mapping and DNA gel blot analysis. The relevant hybridizing bands in each phage were cloned into pBluescript SK+ vectors for further analysis.

Nucleic acid sequencing and analysis

5

25

30

Nucleotide sequencing was carried out manually using the Sequenase Version 2.0 protocols according to the manufacturer's protocol (United States Biochemical, Cleveland, OH) or with an ABI 310 Genetic Analyzer (PE Applied Biosystems, Foster City, CA) using double-stranded plasmid DNA templates prepared utilizing the Qiaprep Spin Plasmid Kit (Qiagen USA, Valencia, CA). The nucleotide and predicted amino acid sequences of the various cDNAs were analyzed using BLAST sequence analysis programs (Altschul *et al.*, 1990; Gish and States, 1993) and protein sequence alignments were carried out using the PILEUP program (Genetics Computer Group Sequence Analysis package, Version 9.0 (GCG, University of Wisconsin, Madison, WI) and the various gene sequences available in the NCBI (National Center for Biotechnology Information, Bethesda, MD) nucleotide and protein sequence database. Manual adjustment of the sequence alignments were carried out as necessary.

RNA isolation and gel blot analysis

Total RNA was extracted from tobacco roots, leaves, and floral parts using Tri-Reagent (Molecular Research Center, USA, Cincinnati, OH) according to the manufacturer's protocol. For RNA gel blot analysis, aliquots (10 µg) of total RNA extracted from the various tissues were fractionated by electrophoresis through a 1.2% agarose-formaldehyde gel and blotted onto Nytran nylon membranes (Schleicher & Schuell, Keene, NH) using 10 X SSC. The transferred RNA was UV cross-linked to the membrane using a UV Stratalinker (Stratagene, La Jolla, CA) and the membranes were prehybridized in 7% SDS, 0.25 M Na<sub>2</sub>HPO<sub>4</sub>, pH 7.2 for 2-4 hours at 65°C. Hybridization was carried out in the same buffer in the presence of <sup>32</sup>P-labeled probes for 16 hr at 65°C. The membranes were washed under high stringency conditions and subject to autoradiography at -80°C for approximately 48 h.

For gel blot analysis, [\alpha-\frac{32}{2}P]-dCTP -labeled probes were prepared by random primed labeling

22

(Random Primed Labeling Kit, Boehringer Mannheim, Indianapolis, IN) using 25-50 ng of a 2.7 kb EcoRI-XhoI fragment derived from PR24 and a 460 bp fragment amplified from the  $\beta$ - subunit of the tobacco mitochondrial ATP synthase gene (atp2) (Boutry and Chua, 1985).

5 Semi-quantitative RT-PCR analysis of NtADC1 and NtADC2 transcript levels.

10

15

20

25

30

Total RNA (2 μg) from roots, leaves, or floral parts was reverse transcribe at 40°C for 1 h in a reaction cocktail containing 200 units of SuperscriptII reverse transcriptase (RNase H-, Gibco BRL, USA), 10 units RNase inhibitor (Perkin Elmer), 200 μm dNTPs and 40 pmol of primer, in total volume of 20μl. For first strand cDNA synthesis, a single primer [5'-AGAAAACATCACCAACT-3'] capable of hybridizing to both the *ADC1 and ADC2* transcripts was used in the reaction. As a control, a primer (5'-GCAACTGTCATCTTATCATCTTC-3') specific for the β-subunit of the tobacco mitochondrial ATP synthase gene *apt2* (Boutry and Chua, 1985) was used in the reverse transcriptase reaction.

Following reverse transcription, the single stranded cDNA products were serially diluted over a concentration range between 1 to 50 ng RNA, and PCR amplification was carried out for 25 cycles of 45 s at 94°C, 1 min at 64°C and 1 min at 72°C in a Genemate thermocycler (ISC Bioexpress, UT). The reaction mixture contained cDNA template, 1 x PCR buffer (Boehringer Mannheim), 100 uM dNTPs, 25 pmol of each forward and reverse primer and 1 unit Taq DNA polymerase. The PCR reactions specific for ADC1 transcripts contained the following primers: ADC1-forward, 5'-CGTAGACGCTACTGTTTC-3' and ADC1-reverse, 5'-TGGACAAC TGTGGAGGCG-3'. Reactions specific for ADC2 transcripts contained primers ADC2-forward, 5'-TGTAGATGCTGCTGTTTT-3', and ADC2-reverse, 5'-TGAACAAC TGCGGAGGCA-3'. Control reactions for normalization of amplification products contained 25 pmol of primers specific for the tobacco apt2 transcripts: atp2 forward, 5'-GTATATGGTCAAATGAATGAGCC-3', and atp2 reverse.int, 5'-GCAGTATTGTAGTGATCCTCTCC-3'. For quantitation purposes, amplification reactions were supplemented with 1µCi <sup>32</sup>P-dCTP. PCR products were separated by electrophoresis through 1.2% agarose gels, the fractionated reaction products transferred onto a Hybond N+ membranes, dried and subject to autoradiography at -70° C. Quantitation was carried out by phosphorimaging using a Molecular Dynamics PhosphorImager. Values were normalized relative to the intensities of the atp2 control band in each lane. The experiment was conducted twice with different total RNA samples, and representative results are presented from one of the two experiments.

23

PCT/US00/12450

#### Results

5

10

15

20

25

30

WO 00/67558

These studies show the structure and expression of individual members of the ADC gene family in tobacco. An  $\alpha$ -  $^{32}$ P-dCTP-labeled 2.7 kb EcoRI-XhoI fragment from PR24 encoding the ADC coding region was used to screen an  $\lambda$  EMBL3 phage genomic library. From a screen of approximately 3 X10<sup>5</sup> phage, seventeen hybridizing phage were recovered, of which five were fully characterized by restriction mapping and DNA gel blot analysis. These phage fell into two groups based on their restriction profile. The relevant hybridizing fragments from the various phage were cloned into pBluescript and their nucleotide sequence determined.

Presented in Figure 7 are the nucleotide and predicted amino acid sequences of NtADC-1 and NtADC-2 genes. Both genes contain a single open reading frame, uninterrupted by introns. The nucleotide and amino acid sequence encoded in NtADC-1 is identical to that of PR24, the full length cDNA isolated from *N. tabacum* cv Burley 21. There are 84 nucleotide differences within the NtADC-1 and NtADC-2 coding regions, resulting in 23 amino acid differences between the ADC1 and ADC2 proteins, respectively. The ADC1 protein is one amino acid shorter in length, missing Val-13.

By comparison to the full-length cDNA, the 5'-untranslated region (UTR) present in NtADC-1 and NtADC-2 are 431 bp and 432 bp long, respectively. The size of the 5'-UTR in the ADC transcripts is considerably larger than the average size of the plant leader sequence (Joshi, 1987). In contrast, the 3'-UTRs present in NtADC-1 and NtADC-2 are relatively short, approximately 84 nucleotides in length. In both gene sequences, a conserved polyadenylation signal (AATAATA) can be recognized 23 nucleotides from the site of polyadenylation site found in the PR24 cDNA.

Pairwise comparison of the *N. tabacum* ADC1 and ADC2 proteins with the ADCs of other plant species showed that the *N. tabacum* proteins are approximately 82% identical to the ADC of its evolutionary progenitor species *N. sylvestris* [Genbank Accession No. AB012873] and 86% identical to the ADC from tomato (*Lycopersicon esculentum*) [31], another member of the Solanaceae family (Figure 2). As might be expected, the *N. tabacum* ADC shares considerably less similarity to ADCs isolated from species more distantly related evolutionarily, such as *Arabidopsis* - 67% identical [32, 33], soybean- 67% identical [34], and oat - 42% identical [35] and is only 29% identical to the enzyme from *Escherichia coli* - [36].

The predicted protein coding regions for the *N. tabacum* ADCs are substantially longer than those reported for the ADC proteins of *N. sylvestris* and *L. esculentum* [31], but are similar in length to those reported in *Arabidopsis*, oat, soybean [32-35] and for the *E. coli* enzyme [36]. The

difference in overall length appears to arise from an apparent nucleotide deletion in the *N. sylvestris* and tomato cDNA sequences relative to the ADC1 and ADC2 predicted sequence and those in other plants. In the nucleotide sequences reported for both the *N. sylvestris* and tomato cDNAs, a guanine residue (position 2295 in the *N. sylvestris* sequence and 1531 in the tomato sequence) is missing [Genbank Accession No. AB012873]. This deletion changes the reading frame and introduces a premature termination to the predicted coding region. Using the sequence information available in the NCBI database, correcting for this error allowed us to extend the predicted C-terminus of the both ADC proteins, yielding the alignment to the *N. tabacum* ADCs and those of other plant ADCs as indicated in Figure 8. We have also included in the alignment shown in Figure 8, the correction at the N-terminus of the predicted tomato ADC protein sequence noted by Pérez-Amado et al. [37], allowing better alignment of all of the higher plant sequences.

Developmental regulation of arginine decarboxylase expression

5

10

15

20

25

30

It has been shown that nicotine formation can be activated in the roots of maturing tobacco plants by topping, that is, removal of the flower head and several young leaves (Akehurst, 1981; Hibi, et al., 1994). Coincident with the activation of nicotine formation, there is an increase in the levels of transcripts encoding ODC, PMT and spermidine synthase (SPS) over the subsequent 24 hr period in wild-type plants (Hibi et al., 1994; Riechers and Timko, 1999). To determine the effects of topping on *ADC* expression in roots, Burley 21 plants were grown in the greenhouse to the bud stage at which point the upper 1/3 of the plant was removed and samples of roots tissues were collected before and at various times post-topping. As shown in Figure 9, *ADC* message abundance increased in the roots of topped Burley 21 plants during the 24 hr period after topping. Low alkaloid (LA) mutants of Burley 21 show a much lower level of ADC expression in their roots, and no induction of ADC transcript accumulation after topping. The lack of ADC induction in the low-alkaloid mutant is consistent with previous studies (Hibi *et al.*, 1994; Riechers and Timko, 1999; Wang, 1999) showing a general inability to activate gene expression leading to increased polyamine formation and alkaloid biosynthesis as a result of the mutation of the *Nic1* and *Nic2* regulatory genes.

NtADC-2 is predominately expressed in roots of wild-type plants.

Due to the high degree of identity between the NtADC-1 and NtADC-2 transcripts (e.g., 95.8% coding regions, 94.4% and 96.4% in 5'- and 3'-UTRs, respectively), it is impossible to distinguish between the two transcripts by RNA gel bot analysis. Therefore, we employed a RT-PCR based

25

PCT/US00/12450

strategy and gene specific oligonucleotide primers. Total RNA was extracted from tobacco roots, leaves and flowers, and single-stranded cDNA synthesized using an oligonucleotide primer capable of hybridizing to both ADC1 and ADC2 transcripts. As an internal control for amplification, a gene specific primer recognizing the *atp2* transcript encoding the β-subunit of the tobacco mitochondrial ATPase was include in the reactions. Under experimental conditions providing amplification in the linear range of the PCR reaction, gene specific forward and reverse primers were used to specifically amplify either ADC1 or ADC2 cDNAs. Test reactions (Figure 10A) using plasmid DNA encoding NtADC1 or NtADC2 as template demonstrated the specificity of the primers. As shown in Figure 10B, the main transcripts detectable in all tissues tested are derived from NtADC-2. Flowers express the highest level of ADC, and leaves lowest. In the flowers, although ADC1 is detectable, far less than ADC2 Roots also express a significant level of ADC.

ADC transcript levels are highest in the roots and floral organs, and low in other plant tissues. The two ADC genes investigated appear to have different modes of regulation, with ADC2 being predominately expressed in the roots and other organs.

At the present time, only limited information is available on the nature of regulatory regions in the promoters of genes encoding enzymes of alkaloid biosynthesis. The availability of cloned genomic fragments encoding ADC allows one to begin mapping regulatory sequences within members of these genes responsible for tissue specific, developmental, and inducible expression.

20

25

15

5

10

#### III. ODC

The present invention features the genes of two members of the nuclear gene family encoding ODC in tobacco *N. tabacum*. As the following experimental discussion shows, the ODC-2 gene is preferentially expressed in roots and floral tissues. Furthermore, the abundance of ODC transcripts in root tissues is affected by topping. Furthermore, analysis of ODC transcript levels in roots of low and high nicotine producing lines shows that ODC expression is under the control of the *Nic1 Nic2* regulatory loci.

## Materials and methods

30

#### Plant growth and tissue preparation

Seeds of *N. tabacum* cv. Xanthi, wild-type and low alkaloid *nic1 nic2* mutant *N. tabacum* cv. Burley 21 were obtained from Dr. G. Collins (University of Kentucky, Lexington). Tobacco plants used for DNA isolation were grown in soil:vermiculite mixture in the greenhouse under natural lighting

5

WO 00/67558 PCT/US00/12450

26

conditions. Plants used for RNA extraction were grown either in Moltan Plus (Moltan Co., Middleton, TN) or hydroponically in a dilute (half-strength) Peters nutrient solution with continuous aeration of the roots under natural lighting conditions in the greenhouse. Topping experiments were conducted by removing the floral meristem, leaves and stem (approximately the upper 1/3 of the plant) from tobacco plants just prior to blooming. Floral parts and other plant tissues were collected from fully matured individuals, frozen in liquid nitrogen, and stored at -80°C until used for RNA preparation (see below).

Screening of genomic libraries and phage characterization

A genomic library constructed in EMBL3 from N. tabacum ev. Xanthi leaf DNA (Clonetech, Inc., 10 Palo Alto, CA) was screened by plaque hybridization (Sambrook et al., 1989) using a <sup>32</sup>Pradiolabeled, 1.6 kb EcoRI-XhoI insert from plasmid PR46 as probe. PR46 encodes a full length ODC cDNA previously isolated by differential screening of plasmid libraries prepared from mRNA isolated from the roots of wild-type Burley 21 plants before and 3-days post-topping (Wang, J., Sheehan, M., Bookman, H. and Timko, M.P., unpublished data). Hybridization was performed at 15 65°C for 16 h in a solution containing 0.25 M Na<sub>2</sub>HPO<sub>4</sub> (pH 7.2) and 7% (w/v) SDS. Following hybridization, the membranes were washed twice in 2 x SSC, 0.1% SDS for 15 min at room temperature, once in 0.2 x SSC, 0.1% SDS for 30 min at 65°C. Hybridizing phage were picked and plaque purified through three subsequent rounds of hybridization screening. Phage DNA was isolated from plaque purified phage using a Qiagen Phage Midi Preparation Kit (Qiagen USA, Valencia, CA) 20 and insert DNA characterized by restriction mapping and DNA gel blot analysis. The relevant hybridizing bands in each phage were cloned into pBluescript SK+ vectors for further analysis.

#### Nucleic acid sequencing and analysis

Nucleotide sequencing was carried out manually using the Sequenase Version 2.0 protocols according to the manufacturer's protocol (United States Biochemical, Cleveland, OH) or with an ABI 310 Genetic Analyzer (PE Applied Biosystems, Foster City, CA) using double-stranded plasmid DNA templates prepared utilizing the Qiaprep Spin Plasmid Kit (Qiagen USA, Valencia, CA). The nucleotide and predicted amino acid sequences of the various cDNAs were analyzed using BLAST sequence analysis programs (Altschul *et al.*, 1990; Gish and States, 1993) and protein sequence alignments were carried out using the PILEUP program (Genetics Computer Group Sequence Analysis package, Version 9.0 (GCG, University of Wisconsin, Madison, WI) and the various gene sequences available in the NCBI (National Center for Biotechnology Information, Bethesda, MD) nucleotide and protein sequence database. Manual adjustment of the sequence alignments were

5

10

15

20

25

30

27

PCT/US00/12450

carried out as necessary.

RNA isolation and gel blot analysis

Total RNA was extracted from tobacco roots, leaves, and floral parts using Tri-Reagent (Molecular Research Center, USA, Cincinnati, OH) according to the manufacturer's protocol. For RNA gel blot analysis, aliquots (10 μg) of total RNA extracted from the various tissues were fractionated by electrophoresis through a 1.2% agarose-formaldehyde gel and blotted onto Nytran nylon membranes (Schleicher & Schuell, Keene, NH) using 10 X SSC. The transferred RNA was UV cross-linked to the membrane using a UV Stratalinker (Stratagene, La Jolla, CA) and the membranes were prehybridized in 7% SDS, 0.25 M Na<sub>2</sub>HPO<sub>4</sub>, pH 7.2 for 2-4 hours at 65°C. Hybridization was carried out in the same buffer in the presence of <sup>32</sup>P-labeled probes for 16 hr at 65°C. The membranes were washed under high stringency conditions and subject to autoradiography at - 80°C for approximately 48 h.

Restriction fragments derived from cDNA clones of interest were separated by agarose gel electrophoresis, the DNA was purified, and quantified by spectrophotometry. [ $^{32}$ P]-dCTP -labeled probes were prepared from 25-50 ng of insert DNA by random primed labeling (Random Primed Labeling Kit, Boehringer Mannheim, Indianapolis, IN). As a control, the blots were also probed with radioactively labeled probes encoding the alkaloid biosynthesis enzyme putrescine N-methyltransferase (PMT) (Riechers and Timko, 1999), a root specific, topping inducible  $\beta$ -glucosidase encoding cDNA (TBG-1) (Riechers, D.E. and Timko, M.P., unpublished data), 26S rRNA (PR31) or 28S rRNA fragments.

#### Genomic DNA isolation and gel blot analysis

Tobacco genomic DNA was prepared from tobacco leaf tissue by the method of Junghans and Metzlaff (1990). Total genomic DNA (15  $\mu$ g) was digested to completion with *EcoRI* or *HindIII*, the digestion products were fractionated by electrophoresis through a 0.8% (w/v) agarose gel, and transferred onto Nytran nylon membrane (Schleicher & Schuell, Keene, NH) in the presence of 0.4 N NaOH (Sambrook *et al.*, 1989). Following transfer, the membrane was rinsed in 2 X SSC, the DNA was UV cross-linked to the membrane, and the membrane was prehybridized and hybridized as described above. Following hybridization and washing, the membranes were subjected to autoradiography at  $-80^{\circ}$ C.

#### Results and discussion

5

10

15

20

25

30

Gel blot analysis of tobacco genomic DNA cut with various restriction enzymes and hybridized with an [α- <sup>32</sup>P]-dCTP-labeled 1.6 kb *Eco*RI-*Xho*I cDNA fragment (PR46) encoding the full-length ODC protein from *N. tabacum* cv Burley 21 (Wang, J., Sheehan, M., Bookman, H. and Timko, M.P., unpublished data) indicated ODC is encoded by small gene family in the *N. tabacum* 

PCT/US00/12450

genome (Fig. 11). Four to five major bands and several minor bands of sufficient size to encode full-length genes are detected in either *EcoRI* or *HindIII* digested tobacco DNA.

To further analyze the structure and regulation of members of the ODC gene family in tobacco, a  $\lambda$  EMBL3 phage genomic library constructed with DNA from N. tabacum cv Xanthi was screened using a [ $\alpha$ -  $^{32}$ P]-labeled probes prepared from PR46 (as described above). From a screen of approximately 3 X10<sup>5</sup> phage, five hybridizing phage were recovered, of which three were fully characterized by restriction mapping and DNA gel blot analysis. Two phage proved to contain identical insert DNA and the third had a unique restriction digestion profile. Following DNA gel blot analysis, the hybridizing fragments were cloned into pBluescript and their nucleotide sequence determined.

The complete *NtODC-2* gene spans two *Sal*I fragments of 2.7 kb and 6.5 kb. The coding region of the gene contains a single1302 bp open reading frame uninterupted by introns (Fig. 12). The nucleotide sequences of *NtoDC-2* is identical within the coding and 5' and 3'- untranslated regions to the PR46 encoded cDNA, with the exception of four nucleotide changes (residues +2, +4, +6 and +8) in the 5'-untranslated region. These nucleotide differences likely reflect changes introduced during the cDNA synthesis reaction.

The predicted amino acid sequence for the *NtODC-2* encoded protein (designated pODC2) (Fig. 13) is identical to the ODC characterized from Burley 21 tobacco encoded by PR46 (Wang, J., Sheehan, M., Bookman, H. and Timko, M.P., unpublished data) and to the partial *N. tabacum* ODC cDNA sequence (PR17) reported by Malik *et al.*, (1996). Comparison of the predicted amino acid sequence for pODC2 with the ODC proteins characterized from two different tobacco cultivars showed that the pODC2 differs by 7 amino acid (98% identity) from the ODC protein characterized from the high alkaloid cultivar, *N. tabacum* cv. SC58 [Genbank Accession No. Y10472.1] and by 7 amino acid (98% identity) from ODC protein from BY-2 cells. The tobacco pODC2 is 89% and 90% identical to the ODCs from tomato (*Lycopersicon esculentum*) and jimsonweed (*Datura stramonium*), respectively, but substantially less similar to ODCs from yeast (35% identity) and humans (32% identity).

The *NtODC-1* gene, contained on an 4.0 kb *XbaI* fragment, encodes a single open reading frame of 141 amino acids encompassing the amino terminal one-half of ODC (Fig. 12). Six amino acid residue changes distinguish the *NtODC-2* and *NtODC-1* encoded proteins over the homologous

5

10

15

20

25

30

region of the proteins. Beginning at amino acid residue 130, the *NtODC-1* encoded protein (pODC1) diverges from pODC2, with a stop codon present after residue 141. Scanning the available nucleotide sequence (> 1 kb) in the 3'-flanking region of the NtODC-1 gene failed to reveal any evidence for ODC homologous protein sequences in any of the three translational reading frames.

29

Interestingly, a comparison of the 5'-flanking sequence of the *NtODC-1* and *NtODC-2* genes revealed that while the *NtODC-2* gene has a clearly recognizable TATA-box properly located at approximately -35 bp from the transcriptional start site, no such regulatory motif is found in the *NtODC-1* gene sequence. Consistent with this observation, RNA gel blot analysis performed using a hybridization probe prepared from *NtOCD-1* immediately downstream of the frame shift, failed to detect any message in various tissues of mature tobacco plants (data not shown). Thus, it appears that *NtODC-2* represents an unexpressed pseudogene in the *N. tabacum* genome.

To determine the spatial pattern of expression of the *NtODC-2* gene, gel blot analysis was carried out using total RNA prepared from roots, stems, young and mature leaves, and various floral parts of Burley 21 tobacco plants. As shown in Fig 14, transcripts encoding ODC were easily detected in the roots, with little or no expression in other tissues except sepals, carpels, and mature stamens.

The formation of nicotine and total leaf alkaloids in tobacco is known to be under the control of at least two independent genetic loci (Legg et al., 1969; Legg and Collins, 1971), designated Nicl and Nic2 (Hibi et al., 1994). Nic1 and Nic2 are semidominant and operate synergistically to control plant alkaloid content, with mutations within these genes resulting in plants with reduced levels of nicotine and total leaf alkaloids (wild-type > nic1 > nic2 > nic1 nic2) (Legg et al., 1969; Legg and Collins, 1971). Although no information is available on the nature of their encoded products, it has been speculated that Nic1 and Nic2 likely encode transcriptional regulators capable of globally interacting with a subset of genes encoding components of polyamine and alkaloid biosynthesis (Hibi et al., 1994). Removal of the flower head and several young leaves (i.e., topping) leads to activation of nicotine formation in the roots of decapitated plants (Akehurst, 1981; Hibi et al., 1994). To determine the effects of topping on NtODC-1 expression in roots, Burley 21 plants were grown in the greenhouse to the bud stage at which point the upper 1/3 of the plant was removed and samples of roots tissues were collected before and at various times post-topping. As shown in Fig 14B, low levels of the ODC transcripts were found in roots prior to topping and message abundance increased approximately 2-fold in the roots of topped Burley 21 plants 4 hr after topping. By 24 hr after topping, ODC transcript levels return to their initial levels. Low alkaloid mutants of Burley 21 subjected to the same treatment show a much lower level of stimulation of ODC transcript accumulation after topping, and the enhanced transcript abundance does not persist beyond 4 hr. By

comparison, transcripts encoding PMT and and a tobacco root-specific  $\beta$ -glucosidase (TBG-1) show patterns of accumulation similar to that observed for ODC transcripts in wild-type plants, but no induction in the low-alkaloid mutant, consistent with previous studies (Hibi *et al.*, 1994; Riechers and Timko, 1999; Wang, 1999).

5

10

15

## IV. SAMS

A single recombinant phage is identified as encoding for SAMS. This  $\lambda$  phage contains an approximately 15kB SalI insert. Restriction mapping and PCR analysis indicates that the insert DNA contains primarily the coding and 3'non-coding portions of the SAMS gene. The nucleotide sequences for the gene encoding SAMS can be found at GenBank Accession Nos. AF27243 (full length SAMS cDNA).

## V. NADH dehydrogenase

A fragment of the cDNA encoding for NADH dehydrogenase in *N. tabacuum* shows high expression in the roots of mature wild-type HP plants compared to low alkaloid mutant LP plants.

## VI. Phosphoribosylanthranilite isomerase (PAI)

The gene encoding for a fragment of phosphoribosylanthranilite isomerase in *N. tabacuum* is a homolog of the *Arabidopsis thaliana* gene encoding PAI, an enzyme involved in tryptophan biosynthesis. This enzyme is involved in the overall formation of aromatic compounds in plants.

## **REFERENCES**

**Akehurst BC.** 1981. The growth, plant structure and genetics. In: Rhind D, Wrigley G, eds., Tobacco, London: Longman Press, 45-95.

25

- **Alabadi D, Carbonell J.** 1998. Expression of ornithine decarboxylase is transiently increased by pollination, 2,4-dichlorophenoyyacetic acid, and gibberellic acid in tomato ovaries. *Plant Physiology* **118**: 323-328.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic alignment search tool. Journal of Molecular Biology 215: 403-410.
  - **Baldwin IT.** 1989. Mechanism of damage-induced alkaloid production in wild tobacco. *Journal of Chemical Ecology* **15**: 1661-1680.

**Baldwin IT, Prestin CA.** 1999. The eco-physiological complexity of plant responses to insect herbivores. *Planta* **208**:137-145.

- Baldwin IT, Schmelz EA, Ohnmeiss TE. 1994. Wound-induced changes in root and shoot jasmonic acid pools correlate with induced nicotine synthesis in *Nicotiana sylvestris* Spegazzini and Comes. *Journal of Chemical Ecology* 20: 2139-2157.
  - Baldwin IT, Schmelz EA, Zhang Z-P. 1996. Effects of octadecanoic metabolites and inhibitors on induced nicotine accumulation in *Nicotiana sylvestris*. *Journal of Chemical Ecology* 22: 61-74.

Baldwin IT, Zhang Z-P, Diab N, Ohnmeiss TE, McCloud ES, Lynds GY, Schmelz EA. 1997. Quantification, correlations, and manipulations of wound-induced changes in jasmonic acid and nicotine in *Nicotiana sylvestris*. *Planta* 201: 397-404.

10

- Bell E.and R.L. Malmberg, Analysis of a cDNA encoding arginine decarboxylase from oat reveals similarity to the *Escherichia coli* arginine decarboxylase and evidence of protein processing. Mol. Gen. Genet., 224 (1990) 431-436.
- **Boutry M. and N.H. Chua**, A nuclear gene encoding the beta subunit of the mitochondrial ATP synthase in *Nicotiana plumbaginifolia*. EMBO J., 4 (1985) 2159-2165.
  - Bracher D, Kutchan TM. 1992. Strictosidine synthase from *Rauvolfa serpentina*: analysis of a gene involved in indole alkaloid biosynthesis. *Archives of Biochemistry and Biophysics* **294**: 717-723.
- 25 Chattopadhyay MK, Ghosh B. 1998. Molecular analysis of polyamine biosynthesis in higher plants. *Current Science* 74, 517-522.
  - **Chou W-M, Kutchan TM.** 1998. Enzymatic oxidations in the biosynthesis of complex alkaloids. *Plant Journal* **15**, 289-300.
  - **De Luca V. and B. St. Pierre.** 2000. The cell and developmental biology of alkaloid biosynthesis. Trends in Plant Science 5: 168-173.
  - Eilbert U. 1998. Induction of alkaloid biosynthesis and accumulation in plants and in vitro cultures

in response to elicitation. In: Roberts MF, Wink M, eds. Alkaloids: Biochemistry, Ecology, and Medicinal Applications. New York: Plenum Press, 219-262.

- Facchini PJ, Penzes-Yost C, Samanani N, Kowalchuk B. 1998. Expression patterns conferred by tyrosine/dihydroxyphenylalanine decarbxylase promoters from opium poppy are conserved in transgenic tobacco. *Plant Physiology* 118: 69-81.
  - Galloway G.L., R.L. Malmberg and R.A. Price, Phylogenetic utility of the nuclear gene arginine decarboxylase: an example from Brassicaceae. Molec. Biol. & Evol., 15 (1998) 1312-1320.
  - Gantet P, Imbault N, Thiersoult M, Doireau P. 1998. Necessity of a functional octadecanoic pathway for indole alkaloid synthesis by *Catharanthus roseus* cell suspensions cultured in an auxinstarved medium. *Plant & Cell Physiology* 39: 220-225.
- Gish W, States DJ. 1993. Identification of protein coding regions by database similarity search.

  Nature (Genetics) 3: 266-272.

10

- Goddijn OJM, de Kam RJ, Zanetti A, Schilperoort A, Hoge JHC. 1992. Auxin rapidly down regulates transcription of the tryptophan decarboxylase gene from *Catharanthus roseus*. *Plant Molecular Biology* 18: 1113-1120.
  - Hashimoto T, Yamada Y. 1994. Alkaloid biogenesis: molecular aspects. Annual Review of Plant Physiology and Plant Molecular Biology 45, 257-285.
- 25 Hashimoto T, Shoji T, Mihara T, Oguri H, Tamaki K, Suzuki K-i, Yamada Y. 1998.
  Intraspecific variability of the tandem repeats in *Nicotiana* putrescine N-methyltransferases. *Plant Molecular Biology* 37: 25-37.
- Hibi N, Higashiguchi S, Hashimoto T, Yamada Y. 1994. Gene expression in tobacco low-nicotine mutants. *Plant Cell* 6: 723-735.
  - Imanishi S, Hashizume K, Nakakita M, Kojima H, Matsubayashi Y, Hashimoto T, Sakagami Y, Yamada Y, Nakamura K. 1998a. Differential induction by methyl jasmonate of genes encoding ornithine decarboxylase and other enzymes involved in nicotine biosynthesis in tobacco cell cultures.

Plant Molecular Biology 38: 1101-1111.

Imanishi S, Hashizume K, Kojima H, Ichihara A, Nakamura K. 1998b. An mRNA of tobacco cell, which is rapidly inducible by methyl jasmonate in the presence of cycloheximide, codes for a putative glycosyltransferase. *Plant & Cell Physiology* 39: 202-211.

**Junghans H, Metzlaff M.** 1990. A simple and rapid method for preparation of total plant DNA. *Biotechniques* 8: 176.

10 Kanegae T, Kajiya H, Amano Y, Hashimoto T, Yamada Y. 1994. Species-dependent expression of the hyoscyamine 6β-hydroxylase gene in the pericycle. *Plant Physiology* **105**: 483-490.

**Kutchan TM.** 1995. Alkaloid biosynthesis - the basis for metabolic engineering of medicinal plants. *Plant Cell* 7, 1059-1070.

**Kutchan TM.** 1998. Molecular genetics of plant alkaloid biosynthesis. In: Cordell GA, ed., The Alkaloids, Chemistry and Biology. San Diego: Academic Press, 295-304.

Legg PD, Collins GB. 1971. Inheritance of percent total alkaloid in *Nicotiana tabacum* L. II Genetic effect of two loci in Burley 21 X LA Burley 21 populations. *Canadian Journal of Genetics and Cytology* 13: 287-291.

Legg PD, Chaplin JF, Collins GB. 1969. Inheritance of percent total alkaloids in *Nicotiana tabacum* L. *Journal of Heredity* 60: 213-217.

25

5

15

Lopes Cardosa MI, Meijer AH, Rueb S, Queiroz Machado J, Memelink J, Hoge JHC. 1997. A promoter region that controls basal and elicitor-inducible expression levels of NADPH: cytochrome P450 reductase (*Cpr*) from *Catharanthus roseus* binds nuclear factor GT-1. *Molecular & General Genetics* 25: 674-681.

30

Malik V, Watson MB, Malmberg RL. 1996. A tobacco ornithine decarboxylase partial cDNA clone. *Journal of Plant Biochemistry & Biotechnology* 5:109-112.

Malmberg RL, Watson MB, Galloway GL, Yu W. 1998. Molecular genetic analysis of plant

PCT/US00/12450 WO 00/67558

polyamines. Critical Reviews in Plant Sciences 17: 199-224.

Michael AJ, Furze JM, Rhodes MJC, Burtin D. 1996. Molecular cloning and functional identification of a plant ornithine decarboxylase. Biochemical Journal 314: 241-248.

5

Mizusaki S, Tanabe Y, Noguchi M, Tamaki E. 1973. Changes in the activities of ornithine decarboxylase, putrescine N-methyltransferase and N-methyl-putrescine oxidase in tobacco roots in relation to nicotine biosynthesis. Plant & Cell Physiology 14: 103-110.

10

Nam K.H., S.H., Lee and J.H. Lee, A cDNA encoding arginine decarboxylase (GenBank U35367) from soybean hypocotyls. Plant Physiol., 110: (1997) 714.

Nam K.H., S.H. Lee and J.H. Lee, Differential expression of ADC mRNA during development and upon acid stress in soybean (Glycine max) hypocotyls. Plant Cell Physiol. 38 (1997) 1156-1166.

15

Ohnmeiss TE, McCloud ES, Lynds GY, Baldwin IT. 1997. Within-plant relationships among wounding, jasmonic acid, and nicotine: implications for defense in Nicotiana sylvestris. New Phytologist 137: 441-452.

20

Pasquali G, Goddijn OJM, de Waal A, Verpoorte R, Schilperoort RA, Hoge JHC, Memelink J. 1992. Coordinated regulation of two indole alkaloid biosynthetic genes from Catharanthus roseus by auxin and elicitors. Plant Molecular Biology 18: 1121-1131.

25

Pérez-Amador MA, Carbonell J. 1995. Arginine decarboxylase and putrescine oxidase in Pisum sativum L. Changes during ovary senescence and early stages of fruit development. Plant Physiology 107: 865-872.

Pérez-Amador MA, Carbonell J, Granell A. 1995. Expression of arginine decarboxylase is induced during early fruit development and in young tissues of Pisum sativum L. Plant Molecular Biology 28: 997-1009.

30

Primikirios, N.I. and K.A. Roubelakis-Angelakis. 1999. Cloning and expression of an arginine decarboxylase cDNA from Vitis vinifera L. cell-suspension cultures. Planta 208:574-582.

20

- **Riechers DE, Timko MP.** 1999. Structure and expression of the gene family encoding putrescine *N*-methyltransferase in *Nicotiana tabacum*: new clues to the evolutionary origin of cultivated tobacco. *Plant Molecular Biology* **41**: 387-401.
- Rostogi R., J. Dulson and S.J. Rothstein, Cloning of tomato (*Lycopersicon esculentum Mill.*) arginine decarboxylase gene and its expression during fruit ripening. Plant Physiol., 103 (1993) 829-834.
- Saito K, Murakoshi I. 1998 Genes in alkaloid metabolism. In: Roberts MF, Wink M, eds.
   Alkaloids: Biochemistry, Ecology, and Medicinal Applications. New York: Plenum Press,147-157.
  - Sambrook J, Fritsch EF, Maniatis T. 1989. *Molecular Cloning: A Laboratory Manual*. Cold Spring Harbor: Cold Spring Harbor Laboratory Press.
- Soyka S and A.G. Heyer. 1999. *Arabidopsis* knockout mutation of *ADC2* gene reveals inducibility by osmotic stress. FEBS Lett 458: 219-223.
  - Stim K.P. and G.N. Bennett, Nucleotide sequence of the *adi* gene, which encoded the biodegradative acid-induced arginine decarboxylase of *Escherichia coli*. J. Bact., 175 (1993) 1221-1234.
  - Suzuki K, Yamada Y, Hashimoto T. 1999. Expression of *Atropa belladonna* putrescine N-methyltransferase gene in root pericycle. *Plant & Cell Physiology* 40: 289-297.
- Wang J. 1999. Characterization of a cDNA (NtcADC1) and two nuclear genes (NtgADC1 and NtgADC2) encoding arginine decarboxylase, a key enzyme in alkaloid and polyamine biosynthesis in tobacco (Nicotiana tabacum L.). M.S. Thesis, University of Virginia, Charlottesville, VA.
- Wang J, Sheehan M, Brookman H, and Timko MP. 2000. Characterization of cDNAs

  Differentially Expressed in Roots of Tobacco (*Nicotiana tabacum* cv Burley 21) During the Early
  Stages of Alkaloid Biosynthesis. *Plant Science* In press
  - Watson M.B. and R.L. Malmberg, Regulation of *Arabidopsis thaliana* (L.) Heynh arginine decarboxylase by potassium deficiency stress. Plant Physiol., 111 (1996) 1077-1083.

Watson M.B., W. Yu, G. Galloway and R.L. Malmberg, Isolation and characterization of a second arginine decarboxylase cDNA from *Arabidopsis* (Ascession No. AF009647 (PGR97-114). Plant Physiol., 114 (1997) 1569.

5

Watson M.B., K. K. Emory, R.M. Piatak and R.L. Malmberg, 1998. Arginine decarboxylase (polyamine synthesis) mutants of *Arabidopsis thaliana* exhibit altered root growth. Plant. J. 13: 231-239.

Waterman PM. 1998. Chemical taxonomy of alkaloids. In: Roberts MF, Wink M, eds., Alkaloids: Biochemistry, Ecology, and Medicinal Applications. New York: Plenum Press, 87-107.

Cizewski-Culotta, V. and Hamer, D.H. 1989. Fine mapping of a mouse metallothionein gene metal response element. Mol. Cell. Biol. 9: 1376-1380.

**Dellaporta, S.L., Wood, J. and Hicks, J.B.** 1983. A plant DNA minipreparation: version II. Plant Mol. Biol. Rep. 1: 19-21.

Foster, R., Izawa, T. and Chua, N.H. 1994. Plant bZIP proteins gather at ACGT elements. FASEB J. 8: 192-200.

**Gerstel, D.U.** 1960. Segregation in new allopolyploids of *Nicotiana*. I. Comparison of 6 x (*N. tabacum* x *tomentosiformis*) and 6 x (*N. tabacum* x *otophora*). Genetics 45: 1723-1734.

25

Gerstel, D.U. 1963. Segregation in new allopolyploids of *Nicotiana*. II. Discordant ratios from individual loci in 6 x (*N. tabacum* x *N. sylvestris*). Genetics 48: 677-689.

Hashimoto, T., Tamaki, K., Suzuki, K. and Yamada, Y. 1998b. Molecular cloning of plant spermidine synthases. Plant Cell Physiol. 39: 73-79.

Hibi, N., Fujita, T., Hatano, M., Hashimoto, T. and Yamada, Y. 1992. Putrescine – methyltransferase in cultured roots of *Hyoscyamus albus*. *n*-Butylamine as a potent inhibitor of the transferase both *in vitro* and *in vivo*. Plant Physiol. 100: 826-835.

Kenton, A., Parokonny, A.S., Gleba, Y.Y. and Bennett, M.D. 1993. Characterization of the *Nicotiana tabacum* L. genome by molecular cytogenetics. Mol. Gen. Genet. 240: 159-169.

- Kim, S.-R., Choi, J.-L., Costa, M.A. and An, G. 1992. Identification of G-box sequence as an essential element for methyl jasmonate response of potato proteinase inhibitor II promoter. Plant Physiol. 99: 627-631.
  - Kinoshita, T., Imamura, J., Nagai, H. and Shimotohno, K. 1992. Quantification of gene expression over a wide range by the polymerase chain reaction. Anal. Biochem. 206: 231-235.

10

20

- Lalanne, E., Mathieu, C., Vedel, F. and De Paepe, R. 1998. Tissue-specific expression of genes encoding isoforms of the mitochondrial ATPase β subunit in *Nicotiana sylvestris*. Plant Mol. Biol. 38: 885-888.
- Legg, P.D. and Collins, G.B. 1971. Inheritance of percent total alkaloids in *Nicotiana tabacum* L.
   II. Genetic effect of two loci in Burley 21 x LA Burley 21 populations. Can. J. Genet. Cytol. 13: 287-291.
  - Leitch, I.J. and Bennett, M.D. 1997. Polyploidy in angiosperms. Trends Plant Sci. 2: 470-476.
  - Li, W.-H. and Graur, D. 1991. Fundamentals of Molecular Evolution. Sinauer Associates, Inc., Sunderland, Massachusetts.
- Mason, H.S., DeWald, D.B. and Mullet, J.E. 1993. Identification of a methyl jasmonateresponsive domain in the soybean *vspB* promoter. Plant Cell 5: 241-251.
  - Menkens, A.E., Schindler, U. and Cashmore, A.R. 1995. The G-box: a ubiquitous regulatory element in plants bound by the GBF family of bZip proteins. Trends Biochem. Sci. 20: 506-510.
- Nakajima, K., Hashimoto, T. and Yamada, Y. 1993. Two tropinone reductases with different stereospecificities are short-chain dehydrogenases evolved from a common ancestor. Proc. Natl. Acad. Sci. USA 90: 9591-9595.
  - Okamura, J.K. and Goldberg, R.B. 1985. Tobacco single-copy DNA is highly homologous to

sequences present in the genomes of its diploid progenitors. Mol. Gen. Genet. 198: 290-298.

Ramsey, J. and Schemske, D.W. 1998. Pathways, mechanisms, and rates of polyploid formation in flowering plants. Annu. Rev. Ecol. Syst. 29: 467-501.

5

Rouster, J., Leah, R., Mundy, J. and Cameron-Mills, V. 1997. Identification of a methyl jasmonate-responsive region in the promoter of a lipoxygenase 1 gene expressed in barley grain. Plant J. 11: 513-523.

Saunders, J.W. and Bush, L.P. 1979. Nicotine biosynthetic enzyme activities in *Nicotiana tabacum* L. genotypes with different alkaloid levels. Plant Physiol. 64: 236-240.

Shinshi, H., Wenzler, H., Neuhaus, J.-M., Felix, G., Hofsteenge, J. and Meins, F. 1988. Evidence for N- and C-terminal processing of a plant defense-related enzyme: Primary structure of tobacco prepro-β-1,3-glucanase. Proc. Natl. Acad. Sci. USA 85: 5541-5545.

Sperisen, C., Ryals, J. and Meins, F. 1991. Comparison of cloned genes provides evidence for intergenomic exchange of DNA in the evolution of a tobacco glucan endo-1,3-β-glucosidase gene family. Proc. Natl. Acad. Sci. USA 88: 1820-1824.

20

15

Staiger, D., Kaulen, H. and Schell, J. 1989. A CACGTG motif of the *Antirrhinum majus* chalcone synthase promoter is recognized by an evolutionary conserved nuclear protein. Proc. Natl. Acad. Sci. USA 86: 6930-6934.

Suzuki, K., Yun, D.-J., Chen, X.-Y., Yamada, Y. and Hashimoto, T. 1999b. An *Atropa* belladonna hyoscyamine 6β-hydroxylase gene is differentially expressed in the root pericycle and anthers. Plant Mol. Biol. 40: 141-152.

Thiele, D.J. 1992. Metal-regulated transcription in eukaryotes. Nucl. Acids Res. 20: 1183-1191.

30

**Thompson, J.D. and Lumaret, R.** 1992. The evolutionary dynamics of polyploid plants: origins,\ establishment and persistence. Trends Ecol. Evol. 7: 302-307.

Vaucheret, H., Vincentz, M., Kronenberger, J., Caboche, M. and Rouze, P. 1989. Molecular

39

cloning and characterisation of the two homeologous genes coding for nitrate reductase in tobacco. Mol. Gen. Genet. 216: 10-15.

Williams, M.E., Foster, R. and Chua, N.H. 1992. Sequences flanking the hexameric G-box core

CACGTG affect the specificity of protein binding. Plant Cell 4: 485-496.

### What is claimed is:

15

20

- 1. An isolated DNA molecule comprising the nucleotide sequence of (SEQ. ID. NO. 2), (SEQ. ID. NO. 5), (SEQ. ID. NO. 8), (SEQ. ID. NO. 11), (SEQ. ID. NO. 13), (SEQ. ID. NO. 15), (SEQ. ID. NO. 15), (SEQ. ID. NO. 16), (SEQ. ID. NO. 17), (SEQ. ID. NO. 18), (SEQ. I
- NO. 18), (SEQ. ID. NO. 21), (SEQ. ID. NO. 23), (SEQ. ID. NO. 25) or (SEQ. ID. NO. 26) or comprising a nucleotide sequence encoding the amino acid sequence encoded by (SEQ ID NO. 3), (SEQ. ID. NO. 6), (SEQ ID. NO. 9), (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) OR (SEQ. ID. NO. 24).
- 2. A vector comprising the isolated DNA molecule of claim 1 operably linked to sequences capable of directing the transcription of a mRNA encoded by said isolated DNA molecule.
  - 3. An isolated DNA molecule comprising a DNA sequence complementary to the nucleotide sequence of claim 1.
  - 4. A vector comprising the isolated DNA molecule of claim 3 operably linked to sequences capable of directing the transcription of a mRNA encoded by said isolated DNA molecule.
  - 5. A cultured transgenic tobacco cell stably transformed with the vector of claim 2.
  - 6. A cultured transgenic tobacco cell stably transformed with the vector of claim 4.
    - 7. A transgenic tobacco plant stably transformed with the vector of claim 2.
- 8. A transgenic tobacco plant stably transformed with the vector of claim 4.
  - 9. The isolated DNA molecule of claim 1, wherein the isolated DNA molecule comprises the nucleotide sequence of (SEQ ID NO:).
- 30 10. A vector comprising the isolated DNA molecule of claim 9 operably linked to sequences capable of directing the transcription of a mRNA encoded by said isolated DNA molecule.
  - 11. An isolated DNA molecule comprising a DNA sequence complementary to the nucleotide sequence of the isolated DNA molecule of claim 9.

- 12. An isolated DNA sequence comprising about a fifteen to about a twenty-five base pair oligonucleotide sequence identical to any consecutive about fifteen to about twenty-five base pair sequence found in (SEQ. ID. NO. 2), (SEQ. ID. NO. 5), (SEQ. ID. NO. 8), (SEQ. ID. NO. 11), (SEQ. ID. NO. 13), (SEQ. ID. NO. 15), (SEQ. ID. NO. 18), (SEQ. ID. NO. 21), (SEQ. ID. NO. 23), (SEQ. ID. NO. 25) or (SEQ. ID. NO. 26).
- 13. A cultured transgenic tobacco cell stably transformed with the vector of claim 10.
- 14.A transgenic tobacco plant stably transformed with the vector of claim 10.

10

15

30

5

- 15. A vector comprising a DNA sequence which encodes an antisense mRNA which is complementary to a fragment of a mRNA encoded by the isolated DNA molecule of claim 1, wherein said sequence is operably linked to sequences capable of directing the transcription of said antisense mRNA in tobacco cells and wherein the expression of said antisense mRNA in tobacco cells is sufficient to provide for reduced nicotine content in tobacco cells which are stably transformed with said vector as compared to untransformed tobacco cells.
- 16. A cultured transgenic tobacco cell stably transformed with the vector of claim 15.
- 17. An isolated and purified protein comprising the amino acid sequence identified in (SEQ ID NO. 3), (SEQ. ID. NO. 6), (SEQ ID. NO. 9), (SEQ. ID. NO. 12), (SEQ. ID. NO. 14), (SEQ. ID. NO. 16), (SEQ. ID. NO. 19), (SEQ. ID. NO. 22) or (SEQ. ID. NO. 24).
- 18. A method for regulating gene expression in a plant comprising functionally linking an alkaloid gene promoter to a nucleic acid encoding a protein, wherein the promoter comprises a nucleic acid sequence selected from the group consisting of the sequences identified in (SEQ ID NO. 1), (SEQ. ID. NO. 4), (SEQ ID. NO. 7), (SEQ. ID. NO. 10), (SEQ. ID. NO. 17), and (SEQ. ID. NO. 20).
  - 19. The method of claim 18, wherein the nucleic acid encoding a protein encodes a protein involved in the biosynthesis of alkaloids in plants.
    - 20. A plant transformed by the method of claim 18.

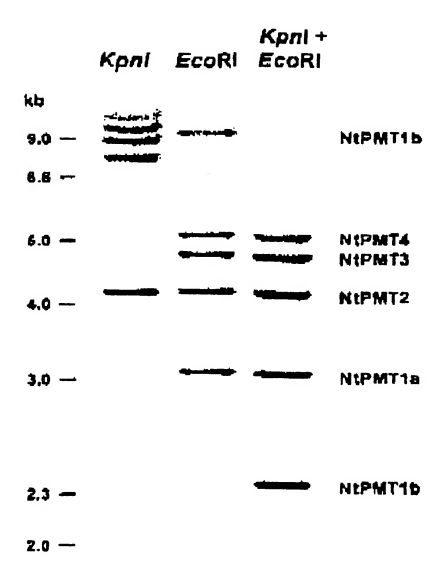


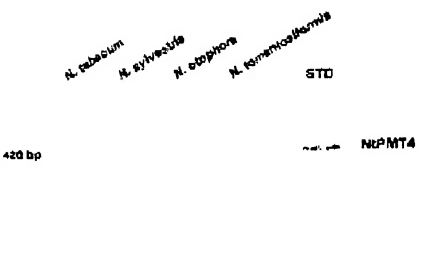
Figure 1

Exon 1

NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	MEVI STNTNGSTI FKNGAI PMNGHQSGTS CHLNGYQNGTSKHQSCHSCHENGY CNGTSKHQSCHSCHSCHSCHSCHSCHSCHSCHSCHSCHSCHSCHSCHS
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	NGHQNG EHQNGHQNGTSE NGHQNG EHQNGHQNGTSE NGHQNG EHRNGHQNGTSE NGHQNG EHRNGHQNGTSE NGHQNG EHRNGHQNGTSE HQNGHQNGTSE
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	QQNGTI SHDNGNE L L GNSNSI K GWF SEFSAL WPG
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP EAFSLKVEKLLFQGKSDYQDVMLFESATYGKVLTLDGAI QHTENGGFP
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	YTEMI VHLPLGSI PNPKKVLI I GGGI GFTLFEMLRYPTI EKI DI VEI D YTEMI VHLPLGSI PNPKKVLI I GGGI GFTLFEMLRYPTI EKI DI VEI D YTEMI VHLPLGSI PNPKKVLI I GGGI GFTLFEMLRYPSI EKI DI VEI D YTEMI VHLPLGSI PNPKKVLI I GGGI GFTLFEMLRYPSI EKI DI VEI D YTEMI VHLPLGSI PNPKKVLI I GGGI GFTLFEMLRYPTI EKI DI VEI D
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	DIVIVIDVISRIKSFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD DIVIVIDVISRIKFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD DIVIVIDVISRIKFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD DIVIVIDVISRIKFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD DIVIVIDVISRIKFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD DIVIVIDVISRIKFPYLAANFINDPRIVITLIVLIGDIGAAFIVKAAQAGYYDAIII VD
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	Exon 6  SISDIPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I SSDPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I SSDPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I SSDPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I SSDPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I SSDPII GPAKDL FERPIF FEAVAKAL RPGGVVCIT QAESII WL HMHI I KQI I
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	Exon 7  ANCRQVFKGSVNYAWTTVPTYPTGVI GYMLCSTEGPEVDFKNPSNPI D ANCRQVFKGSVNYAWTTVPTYPTGVI GYMLCSTEGPEVDFKNPVNPI D ANCRQVFKGSVNYAWTTVPTYPTGVI GYMLCSTEGPEVDFKNPVNPI D ANCRQVFKGSVNYAWTTVPTYPTGVI GYMLCSTEGPEVDFKNPVNPI D ANCRQVFKGSVNYAWTTPTPTTGVI GYMLCSTEGPEVDFKNPVNPI D ANCRQVFKGSVNYAWTTVPTYPTGVI GYMLCSTEGPEWDFKNPVNPI D Exon 8
NtPMT4 NtPMT3 NtPMT1a NtA411 NtPMT2	KETTTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES KETTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES KETTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES KETTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES KETTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES KETTOVKSKLAPLKFYNSDIJHKAAFILPSFARSMIES

Frqure 2

3 / 21



31 Ó ÞA	<b>8</b> In	-		Septe of all	NIPMTS
290 bp NEPMT18			-	المرجعين	NEPSETTS

220 op NEPMTZ

Frque 3

					-				
NtPMT3				gc	tgtacaaaag	gatgtctcaa	atcatttgga	atattaattc	-1029
NtPMT2 NtPMT4								ctgagttg	-1039
Nt PMT1a									
NtPMT3	+							DOX	
NtPMT2	acaagaacaa	aagaaatacc ttcctggtga	atcagatgga	tgaagataat	agaggtagat	aaaattatga	gatcattaaa	catcttaaac	-949
NtPMT4							accasageag		-333
NtPMT1a									
NtPMT3	ctatecetat	ttggaagagt	ataatataaa	2021000100	anaant	*****	******		0.00
NtPMT2	gactgtgcga	ttggaagagt gttgcagaaa	caattgaagg	gtcatttgtg	gaatttgggg	ccatttcaaa	ggaaaaagaa	aagatgactt	-879
NtPMT4									0.5
NtPMT1a									
NtPMT3	acaaattggg	aagcagggat	tgagggattc	tcagagatga	agagggaggc	tttatcataa	ctttttcaat	acctataatc	-789
NtPMT2	agcattaata	aatcaaatta	aaataaggct	tagcgttaaa	atcaaaggaa	atggcaagcc	tggctcctgg	agcaatgctt	-799
NtPMT4 NtPMTla									
NCFMIIA									
NtPMT3	tataataaca	tcagtgaagc	agaattgaaa	gccatcaagt	atgggtgtga	atggtgcaaa	tacaaaggaa	tatcaaactt	-709
NtPMT2	ctgaggacag	tagtaaaaac	aatatcagac	aaaaagtaaa	gttgtattat	ttagcttgag	gataaagtat	gtcattagtt	-719
NtPMT4 NtPMT1a									
Nt PMT3	cattgtggaa	actgactcga	ggatgatcta	tgacatacta	cagaccaaaa	#tc#aagcaa	caacaagttg	aaacaagaga	-629
NtPMT2 NtPMT4	ttgtgagaga	tttggtgtcc	tctacaatga	ttgttgaagt	ccctatttat	#gc#atacac	aggaaacaaa	atcctaggat	-639
NtPMT1a						*at*caatgg	agaaggaaaa		-625
					7	##	agaaggaaaa	ta to to to tag t	023
M+ DMT 2				Box			0000	**	
NtPMT3 NtPMT2		aatggagatt ttaaatgaca							
NtPMT4							acycomman.		-339
NtPMTla	gtaaacacaa	gtgaatgaag	agaagccaaa	ataatctcta	tcattcaagc	cttaggtgga	gatta <b>na</b> aaa	at#atttact	-545
						PAL			
NtPMT3	tggtttgcta	aagäggccac	cagagetaac	gaaggtätcä	Etcattacaga	ttttagacag	gtatcaaaag	cggccaaggg	-469
Nt PMT2	gactatttat	ttaätattga	ggaatatttt	ttatta@at@	*tatctggtg	acaagcattc	gtttgcttcc	gttgattacg	-479
NtPMT4 NtPMT1a		agt <sup>®</sup> ataggt							
NtPMT3	ccctttcktc	atggatatg§	ggcaggtccc	ttattttaga	attaga <b>k</b> atg	aaaaa§ctaa	ttttttttg	taagttaatt	-389
NtPMT2 NtPMT4	ttgattt	gatctactc	ataccaaccg	aagccgttgt	ccttgagctt	cgctt&catt	taattcatct	tccgtatgcc	-399
NtPMT1a	acccacagaa	ggaagatca§	aaaatacatg	actttcagat	gacttcttgg	agcttEattt	ttaaagagtg	gctaggacc	-385
NtPMT3 NtPMT2	ctgtgtatag	tgagaggaaa	tcgtctaata	tgtattttt	cccatagact	cttostctcc	ttaggtaaaa	aggtagctc	-309
NtPMT4	tctgattcca	cangtdatgc cangtdatgc	acccattcas	ttatttaatu	gamaccamos gamaccamos	transctata	caaatoggtac	atcattcgte	-319
NtPMT1a		gtgcttggtca							
NtPMT3	-×		333_35	**********			_87 39 .86	. 2000	
NtPMT2	aaatecteta	ttatättccc cttggatata	aasaasttta	ENGCOSOGACT	Raacamataca	gacatggtat	gggtccagcs	taaanaaata	-229
NtPMT4	aaatactatc	cttggataaa	aadaattttg	###ta#ggagt	- Baacagatgc	gaagtaagaa	apcagacgac	taaagaaaat	-239
NtPMT1a	acaaaagkaa	gaggaaggaaa	ggagacagaa	gaggäaatag	ättt <u>g</u> ggggg	<b>\$</b> 99 <b>\$</b> 999 <b>\$</b> 99	gtttcagaat	caaagaasat	-225
NtPMT3	caccäcännä	patagatacc	****aataatta	otttatt%tt	#3333333	22200	**********	#_88_####	-153
NtPMT2	ttt#####a	gagagagazz	tgaamacaca	gatgtactaa	taaaattagg	gtactactat	actaataatt	SCHOOL SEC	-159
NtPMT4	tttassaaag	dadededass	tgaguacaca	dacgtactaa	taaaattagg	gtactactat	acteateatt	ggagagagac	-159
NCPMTIA	rrressaatg	fadededars	EGSGRACSCS.	mara Escret	C#####		acceateatt	gcadcuagas	-156
		cc	AAT-Box		G-Bo	k PAL			
NtPMT3	aaääc≹tätä	1111.001.1E2	aaaatata	cagtera	acca geacq	ttgraatgat	tittiaau	-totattata	-84
NtPMT2 NtPMT4	taaastcata	tittagtiks	aaaat#tctc	gggeagteus	accadacaco	ttotastgat	tttttaae	-tetattata	-82
	aaactt-ata	tretagete	aaaatut	caetet#	acceptance	110121112	rttttaau	tvaisttata	-84
_						· · · · · · · · · · · · · · · · · · ·	outermound and	-xx=-exxenses	- •
Me numa	))))))))	re : :::::::::::::::::::::::::::::::::::	2222	TATA	l-Box	_2222222222	202000000000000000000000000000000000000	202022200000000000000000000000000000000	
NtPMT3 NtPMT2	togentiess	ceetecacie	ctemmates	properties t	razargeeta	gatarotti	regggagtat	acatcaagct	-4 -4
NtPMT4	togenitoca	ccctc acto	etetyratee	aughtetara	tazatucet-	-steresca	troppageor	acagggaact	-4
Nt PMT1a	togeg toog	ccctcdectc	ctcggratcc	seettotet	taantgcet-	-agatgttta	trassastar	acagement.	-4
	+1							\	
NtPMT3	ttcecaaaat	eca		Beccatasta	ELECCIONE.	ECCHOEREN-	tranterape	MET	+60
NtPMT2	ttcetaaægt	eca ecaaatcgta ecaaatcata eca	atacttgttg	Beacataste	ctttctcttc	tcceetttgt	ttagtttaet	ittgeeeatg	+77
NtPMT4	ttcetaaagt	ecaaatcata	atacttgttg	eeacataste	ctttctcttc	tcceetttgt	ttagttteet	ittg###ATG	+77
MCEMITA	****Adawa*	*****		##CDADABAGE	MINISTERIC	*cementals	***********	LCCCHOOAIG	+60

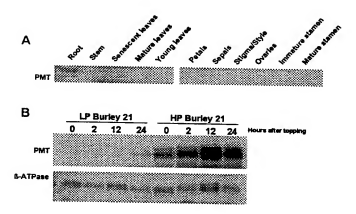
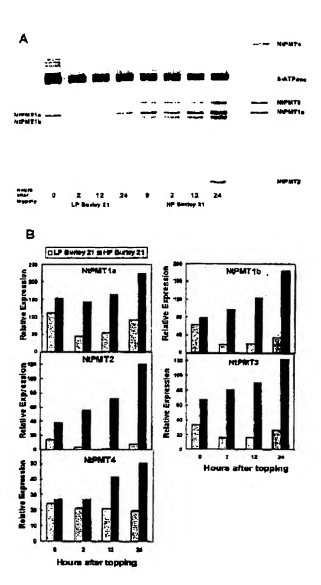


Figure 5

6 / 21



trquelp

NEADCI	
ttcacgttctcttctcaattcccataaaagaaacccttc	egttag. <sup>319</sup>
gtttccgtcctattttctcttcttctacgcttc 78 NtADC2tcact80 NtADC1	NtADC1 agccccatctattacaaccattgggcaaaaacatca ttaaatctgtacaaagcaaacccttaatttagtttaa ttttct 398 NtADC2
ctcttctgatatcaatatctgtatggtgtttttcttg ttcgaattttagatttgttttgcctttaatacctgta acctta 158 NtADC2	at
ataaa	1 M P A L G C C V D A T - V S P P NtADC1
NtADC1 taattototgtttaaaccaaaaacttagcttottotg aagtcagggtggggatatttggatcgtgtaagagtgt gttaga 238	gtattctttgattctttaacagaagaagaagagATGCCGGCCCTAGGTTGTTGCGTAGACGCTACTGTTTCCCCTCC 475 NtADC2
NtADC2t	atTGGTT 479
NtADC1 aggtgattatcttttgattcagttccttttttgcttc	M * * * * * * * * * * * * * * * * * * *
ttttgaggggtagccggggcctcggcctcggcggt tttaat 318 NtADC2	16 L G Y A F S R D S S L P A P E F F T S G V P P T N S A NtADC1
4	$CTCGGCT\Delta TGCCTTCTCTCGGG\Delta T\Delta GCTCTCTTCTCCCG$

PCT/US00/12450

		ننAG		TTT.	ACC	TCC	GGC	GTF	ICC.	CC	TA	CA.	AA		ADC	_										
		55	5														т.								.c.	
Νt	ADC	:2															Γ		Γ							
																. 7	719									
					G.,						G.				7	0 *	k 1	k 1	k .	*	* :	* 1	k 1	. ,	k y	•
	.T.	55	9											*	*	*	*	*	*	*	*	*	*	*	*	*
	1	.7	*	S	*	*	*	*	*	*	*		*	*	*	*	*									
*	*	*	*	*	*	*	Α	*	*	*		*	*													
*	*	*	*												9	6	K	ĸ	A	s	D	P	к	N	S	G
														G	L	G					L					
	4	.3	A	G	s	I	G	s	P	D	1	L	s			L		_	_	_		-	•	••	•	-
s	Α	L				D							Y	Nt	ADC	1.										
F	s	v					_		_				_			_	ייטיי	רככ	TAC	רת:	AAA	ידעע	רכשכ	200	יממני	יידיר
Nt	ADC	:1																			rtg:					
CC	GCC	GGT	TCC	ATT	GGG	тст	CCG	GAT	CTC	TC	СТ	77	GC			A 7						(			.102	110
		'ACG													ADC		, , ,									
		T 6																						T		T.
	ADC													• •												
		.c.							d.		c			• •		 		• • •	• • •	• • •	• • •	• • •	• • • •	• • •	• • •	• •
														• •		7	*	*	*						_	_
		. 6		• • •	• • •	• • •	• • •	• • •	• • •	• •	• •	• •	• •			, , •						٠.	٠.			^_
• •		4		A	*	*	*	•							Ĵ			-	-	•	_	•	•	•	•	*
*	*	*	*	*	*	*	*	*					Ĵ	-	-	•	_									
*	*	*													12		N	ъ	т.	F	s	т.	_	c	7	_
														D		_					G					
	6	9 N	S	N	G	D	т	9	, ,	,	Ð	ъ			G		٧	**	٥	Q	•	-	G	А	п	ī
н	G	T											т.	~	ADC	-										
		v		-	_	-	••	×	-	_	•	_				_	ייייי	מבים ז	ייייי	ירית	GCA/	ייי מי		بالملاملات	רכי איז	7
_	ADC	•	•																		rat(					
		TAA	CGG	AGA	ייבי	רידירי	ርርጥ	רכי	:אכיר	יאמי	<u>አ</u> ጥረ	.ت	מיד			T 8			-76	<b>3</b> GC.	TVT(	الحال	3000	.nc.	MCC	.AA
		ACA													ADC	_	, , ,									
		T 7			CAC	CAG		-> T T	GAC	1	<u>.</u> .		~~													
-																										

124		.G.													17	7	*	*	*	*	*	*	*	*	*	*
* * * * * * * * * * * * * * * * * * *			. 8	79										*	*	*	*	*	K	*	*	*	*	*	*	*
203   F   K   D   A   E   Y   I   S   L   A   L   V   A   R   K   L   M   L   N   T   V   I		12	24	*	*	*	*	*	*	*	*	*	*	*	*	*	*									
L V A R K L M L N T V I V  149 Y P V K C N Q D R F  V V E D I V K F G S S F R  NtADC1  TATCCCGTGAAATGCAATCAAGACAGGTTCGTGGTGG AAGATATTGTCAAATTCGGGTCGTCGTCGGTTCGG GTTGGA 955  NtADC2	*	*	*	*	*	*	*	*	*	*	*	*	*													
L V A R K L M L N T V I V  149 Y P V K C N Q D R F  V V E D I V K F G S S F R  NtADC1  TATCCCGTGAAATGCAATCAAGACAGGTTCGTGGTGG AAGATATTGTCAAATTCGGGTCGTCGTCGGTTCGG GTTGGA 955  NtADC2	*	*	*												20	3	F	ĸ	D	Α	E	Y	т	S	т.	Δ
149 Y P V K C N Q D R F														Τ.	v	_ A	R	к			_	_			_	
V V E D I V K F G S S F R NtADC1  F G L E		14	19 Y	P	v	к	С	N	0	D	R	F	7	T.	E	0			_	• • •	_		•	•	-	٠
## Company of the com	v	V	E	D	I	v	ĸ	F	~			_	R	N+	ADC	~										
TGCAAGAAAGCTCATGTTAAACACTGTAATTGTTCTT	F	G	L	E	_			_	•	-	_	-					CAC	יניריז	יכאכ	יתאר	יייי מי	יייירים	ירידים	יכיכים	المادلات	-C-T-
TATCCCGTGAAATGCAATCAGACAGGTTCGTGGTGG AAGATATTGTCAAATTCGGGTCGTCATTCCGGTTCGG GTTGGA 955 NtADC2	Nt.	ADC	21	_																						
AAGATATTGTCAAATTCGGGTCGTCATTCCGGTTCGG  GTTGGA 955  NtADC2				מעם	מדמ	יב בי	TCA	מבאמ	מסמר	ייייב	مريب	ССТ	cc						.101		Inch	.CIG	TMM	.110	2 T T (	-11
STTGGA 955   S																		,								
NtADC2					rurura.	110	GGG.	LCG.	LCM.	110	ىقى	110	.GG	14 C												
				23										• •												
204 * * * * * * * * * * * * * * * * * * *														• •					• • •	• • •	• • •	• • •	• • •		.G	• • •
														• •		-	.119	,								
150					• • •	• • •	• • •	(	Z.,	• • •		• • •	• •		20	4	*	*	*	*	*	*	*	*	*	*
* * * * * * * * * * * * * * * * * * *	• •			59										*	*	*	*	*	*	*	*	*	*	*	*	*
* * * * * * * * * * * * * * * * * * *		15	50 <b>*</b>	*	*	*	*	*	*	*	*	*		*	*	*										
S R K M A V R P V I G L R  176 A G S K P E L L L A A K L R  M S C L C R G S A E G L L  V C N G  NtADC1  AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC TGTCTCTGCAGGGGGCAGTGCTGAGGGCCTTCTCGTTT  GCAATG 1035	*	*	*	*	*	*	*	*	*	*	P	*	*													
176 A G S K P E L L L A A K L R  M S C L C R G S A E G L L  V C N G  MtADC1  AGATGGCTGTTCGCCCGTAATTGGACTTCGGCTAA  AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC  TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT  GCAATG 1035	*	*	*	*											22	9 E	: E	E	I	ı E	L	v	I	Γ	) :	I
M S C L C R G S A E G L L V C N G GAGGAGGAGCTTGACCTTGTGATTGATATAAGCCGTA NtADC1 AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT GCAATG 1035  NtADC1 AGATGGCTGTTGGCCCTGATTGGACTTCGGGCTAA AGATGGCTGTTCGGCCCGTAATTGGACTTCGGGCTAA AGATGGCTGTTCGGCCAGACTTCTCGTTT NtADC2														s	R	K	M	Α	V	R	P	V	I	G	L	R
V C N G GAGGAGGAGCTTGACCTTGTGATTAAAGCCGTA NtADC1 AGATGGCTGTTCGGCCCGTAATTGGACTTCGGGCTAA AGATGGCTGTTCGGCCCGAATTGGACTTCGGGCTAA AGATGGCTGTTCGGCCCGTAATTGGACTTCGGCCTAA GCTCAG 1195 NtADC2 GCAATG 1035		17	6 .	A (	G :	s :	K :	P 1	E ]	<b>L</b> 1	ւ :	L	A	A	K	L	R									
NtADC1 AGATGGCTGTTCGGCCCGTAATTGACTTCGGCCTAA AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT GCAATG 1035  AGATGGCTGTTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTTCGGCCTAATTGGACTTCGGCCTAA AGATGGCTGTTCGGCCTAATTGGACTTCGGCCTAA ACATGGCTGTTCGGCCTAATTGGACTTCGGCCTAA ACATGGCTGTTCGGCCCGTAATTGGACTTCGGCCTAA ACATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA ACATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA ACATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCAGGCCTCCGGTT ACATGGCTGTCGGCCAGGCCTCCGGTT ACATGGCCGGCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCAGGCCCTCCGGTT ACATGGCCGGCCTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCAGGCCCTCCGGTT ACATGGCCGGCCTCCGGTT ACATGGCCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGTCGGCCCGTAATTGGACTTCGGCCTAA AGATGGCTGCGGCCTTCTCGGTT ACATGGCCGGCCTAATTGGACTTCGGCCTAA AGATGGCTGTAATTGGACTTCGGCCTAA AGATGGCTGGCCGGCCTTCTCGGTT ACATGGCCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAA AGATGGCTGGCCGGCCTTCTCGGTT ACATGGCCGGCCTTCTCGGTT ACATGGCCGGCCTTCTCGGTT ACATGGCCGGCCTTCTCGGTT ACATGGCCGGCCGTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTTCGGCCTAATTGGACTGACT	M	S	C	L	C	R	G	S	Α	E	G	L	L	Nt	ADC	1										
NtADC1 AGATGGCTGTTCGGCCCGTAATTGGACTTCGGGCTAA AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC GCTCAG 1195 TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT NtADC2 GCAATG 1035	v	С	N	G										GA	GGA	GGA	GCI	TGA	CCI	TGI	GAT	TGA	TAT	AAC	CCC	TA
AGCTGGGTCTAAACCCGAGCTCCTGTTAGCCATGAGC GCTCAG 1195 TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT NtADC2 GCAATG 1035	Nt	ADC	:1																							
TGTCTCTGCAGGGGCAGTGCTGAGGGCCTTCTCGTTT NLADC2 GCAATG 1035	AG	CTG	GGT	CTA	AAC	CCG	AGC:	rcc:	rgT:	rag	CCA'	TGA	GC													
GCAATG 1035A														Νt	ADC	2										
																_									,	۸.
C																				• • •	• • •	• • •	• • •	• • •	• •	• • •
														• •												
1039 * H * * * * * * * * * * * * * *					• • •	• • •	• • •	• • •	• • • •	• • •	• • •	• • •	• •			J ^	<u>.</u>		Ţ,	ı.		_ ~				·
* * * *	• •	• • •		<b>433</b>										_		_	Ĵ	-	•	^	^	~	*	*	*	*

10 / 21

256 T K H S G H F G S T SGEKGKFGLTTTQ NtADC1 I V R V ACGGCGTTGCTTGCTGATGGTGTGAGGCTGCTC NtADC1 AGATTTATTGTGAATTAATCCGTCTTGGTGCGGGTAT GACCAAGCATTCAGGCCATTTTGGATCCACTTCTGGA GAAGTT 1435 GAAAAAGGTAAGTTTGGGCTTACAACGACCCAAATTG NtADC2 TTCGTG 1275 NtADC2 ..... 1439 310 \* G \* ..... 1279 257 \* 336 I D T G G G L G I D 283 V K K L E E S G M L YDGTKSCDSDVSV D C L Q L L H F H I G S Q GYGI I P S NtADC1 NtADC1 CATTGATACTGGAGGTGGGCTCGGAATTGATTATGAT TAGTGAAGAAGCTGGAAGAATCCGGAATGCTGGATTG GGTACTAAATCATGTGATTCAGATGTCTCTGTTGGCT CCTTCAGTTGCTGCATTTTCACATTGGATCTCAGATC ATGGCA 1515 CCTTCA 1355 NtADC2 NtADC2 T............ ..... 1519 ....T 1359 337 \* \* 284 \*

	36	3	Q	E	Y	A	S	T	V	v	Q	Α	Nt.	MC:	1										
V	Q	Y	v	C	D	R	K	G	V	K	Н	P	TTC	TT	CAC	ATC	TGT	CTT	CTG	GTG	GCC	TCC	'AAT	CCA	TG
V	I	С											GCG	GAG	GAC	GCT	CAA:	ГGА	AGA	TGC	CCI	TGC	TGA	TTA	.CC
Nt.	ADC:	1											GCA	AT:	r 1	755									
TT	CAA	GAA	TAC	:GCC	TCC	ACA	GTT	GTC	CAG	GCG	GTT	'CA	NtA	DC:	2										
					GTA																				
	TTG																								• •
	ADC:														. 1		• • • •	- • •	• • •	•••	• • •		• • •	• • •	• •
			т	•		C		т	,				• • •	41		, *	* ;	*	*	*	*	+			4
						-		–							<b>,</b>										
	c				• • •		• 1 •	• • •	.л.	• • •	• • •	• •			··	 -				•	-	-	•	-	_
• •	36.	-	פפכ. •	<u>.</u>		_			*				•	•	•	•									
_	36	4	_	_	Î.	_	_	A		<b>*</b>		*			_	_	~	_	_		_	_	_		_
×	.a.	<del>-</del>	*	*	*	*	*	*	*	*	*	*	_	44	_	S	A	A	A	V	R	_	E	Y	E
*	*	*											T	C	•	L	Y	s	D	Q	L	K	Q	R	С
				_		_	_	_				_	V	D	Q										
		9 S	_	-	_					_			NtA												
H	S	I	L	I	F	E	A	v	S	A	S	S											GAG		
H	s	С	S															ATC	AGI	'TGA	AAC	'AGA	GAT	GTG	TG
	ADC												GA?	CA	G 1	835									
AG	CGA	AAG	TGG	CAC	GGC	AAT	TGT	TTC	TCA	TCA	CTC	AA.	Nt.	ADC:	2										
TT	CTG.	ATT	'TTC	GAA	.GCC	GTG	TCT	GCI	TCI	'AG'I	CAC	TC											· · ·	A	
AT	GTT	C 1	675	;																					
Nt	ADC.	2													. 1	839									
														44	4	*	*	*	*	*	*	*	*	*	*
													*	*	*	*	*	*	*	*	*	*	*	*	*
		. 1	679	)									*	*	*										
	39	0 *	*	. *	*	*	*	*	*	*	*														
*	*	*	*	*	*	*	*	*	*	*	*	*		46	9 F	K	E	G	S	·	, (	3 I	E	Н	[
*	*	*	*										L	Α	Α	v	D	s	I	С	D	F	v	S	ĸ
													А	M	G	Α									
	41	6	s	s	н	L	s	S	G	G	L	0	Nt.	ADC	1										
s	М	A	E	T	L	_ N	E	_ D	A	L	_ A	*D				AGG	GTC	СТТ	'GGC	י בידי	тĢz	ACZ	TCT	TGC	יידיבי
v			т.	_	_		_	_		~	••	•													- 0

~m	~~~	~~ m		3 mc	mam	~ n m	<b></b>	am 3	mai				37L 7												
					TGT	GAI	111	GIA	TCA	AAA	1 کی	AT.		ADC2				_		~					
	GGG		9 <b>T</b> 2																	c					• •
Nt.	ADC:	2											• • •	• • •			• • •	• • •	• • •		• • •	.A.	• • •	• • •	• •
							• • •					• •			. 20	079									
														524	Į.	*	*	*	*	*	*	*	*	*	*
		. 1	919										*	*	*	*	*	*	*	*	*	*	*	*	*
	47	0 *	*	*	*	*	*	*	. ,	* 1	* 1	+	*	*	*										
*	*	*	*	*	*	*	*	*	*	*	*	*													
*	*	*	*											549	9 G	к	v	D	K	F	I	G	G	E	
													S	S	L	0	L	н	E	L		s	N	G	D
	49	6	A	n	P	I	D	т	Y	н	v	N	G	G	_	Ϋ́	_		_	_	_	_		_	-
L		-	F	_	s			_			A		-	ADC:	_	-									
	_		ı T	1	3	1	P	ט	r	W	A	ŗ				יסאי	TI A A	~mm	(1) III	maa	maa	003	3 m.a		-
G	-		r																	TGG					
	ADC	_															GAA	TTG	GGA	AGT.	AAT	GGC	GAT	GGT	GG
					GCA									GT?		155									
AT	TTT	CAC	TTC	raa:	TCC	TGA	TTT	TTC	GG	CCT:	rtg	STC	Nt	ADC:	2										
AΑ	TTG	T 1	995	,																					
Νt	ADC	2												.c.	A										
			G	;										.T.	. 2	159									
														55	<b>)</b> *	*	*	*	*	*	*	*	*	*	:
		. 1	999	)									*	*	*	P	*	*	*	*	*	*	*	*	*
	49	7	*	*	*	ν	*	*	*	*	*	*	*	*	*	*									
*	*	*	*	*	*	*	*	*	*	*	*	*													
*	*	*	*											570	٠ .	Y	L	c	M	다	L	G	G	Α	Y
													E	•	_	_			L		N	_	F	G	G
		_	_	_	**	_	_		_		_	_	P.		V	_	G	G	П	п	1/4	ъ	F	G	G
	52	_	P		V					L	_	E	-	_	-	٧									
K	_	A	٧	R	G	T	L	S	D	L	T	C		ADC:	_										
D	S	D																		GTG					
Nt	ADC	1															ACT	CCA	CAA	CCT	GTT	TGG	TGG	ACC	AA:
TI	'CCG	ATI	GTI	CCI	ATA	CAC	CGT	TT	\GA'	TGA	AAA	GCC	GC	GTG(	G 2	235									
TG	CAG	TAP	GGG	GAZ	IAT!	TAT	'CGG	AC:	CTG.	ACT'	TGT	GAC	Nt	ADC:	2										
AG	TGA	T 2	075	5																					

		• • •	• • •											63	0 *	•		,	* 1	* *	4	* *	*	*	!
.T	c	. 2	239										*	*	*	*	*	*	*	*	*	*	*	*	*
	57	7	*	*	*	* :	*	*	*	*	*	*	*	*	D	*									
*	*	*	*	*	*	*	*	*	*	*	*	*													
*	*	*	*											65	6	ĸ	G	τ.	A	Т	Α	S	L	Δ	S
													S	L									_ Y		_
	60	3	R	v	v	0	s	D	S	Δ	н	s		P		Š	_	-	**		•••	•	•		٠
F	A	_			s				P		C	Δ		ADC		_									
_	v	Τ.	_		_	•	-	_	•	~	•	••			_	יידיכיני	ברכז	ידידי	ימיזי	יישיטי	רידיכים	cci	\GCA	CCT	אידיי
_	ADC	_																					TGI		
		_	GTG	מאמי	אמרי	ייי מב	A C C C	ممت	מא מי	אככי	ጥጥር			GCA				11.77	·	GCC	- 1 12	3CC 1	. 1 G 1	ناتات	.پ
			GCT											GCA ADC		4/5	•								
		C 2		CCG.	100	- 1 G	المال	CGI	CCI	تاعی	CGG	3AC	_				me		70					_	
	ADC		313																						i
																		• • •	• • •	• • •	• • • •	• • •	• • •	• • •	• •
													• •	T	-		,	_			_				
			• • •	• • •	• • •	• • •	• • •	• • •	т.	• • •	т.	T.			7	* .	* .	* .	* .	٧.	E	*	*	*	*
• •		. 2							_					V		*	*	*	*	*	*	*	*	*	*
	60	4	*	*	*	*	*	*	*	*	*	*	*	*	S	*									
*	*	*	T	*	*	*	*	*	*	*	*	*													
*	*	*												68	_								Α		
													G	G	Y	N	Y	Y	Y	S	D	E	N	Α	Α
			A								M	1	D	S	Α										
F	E	T	L	K	H	R	A	E	$\mathbf{E}$	F	L	E	Nt	ADC	1										
Q	E	E	D										CT	TGC	TGC	TTC	CAC	rgc2	AGT:	rac:	rgc1	CAAC	CAAC	:GGT	'GG
Nt	ADC	1											CT	ATA	ACI	'AC'	CTAT	CAC	AGT	SATO	SAG	ATO	CAG	CAC	:AT
CG	AGC	GAT	GCA	GCA(	CGA	GCC(	CGA	GCT	CAT	GTT	CGA	\GA	TC	TGC	т 2	555	5								
CT	CTC.	AAG	CAC	CGT	GCG	GAG	GAA'	TTC	TTG	GAA	CAA	\GA	Nt	ADC	2										
AG	AAG.	A 2	395												С.,		. <b>.</b>		r.c		A.		т	·	
Nt	ADC.	2													т.										
															. 2	559	•								
														68		*	R	*	*	*	A	*	D	*	*
	т.	. 2	399						•	•			*	*	*	*	*	*	*	*	*	*	*	*	*

14 / 21

	709	т	G	E	D	E	I	W	s	Y	С
T		***							_	-	•
<b></b>											
	DC1										
		GAG									
GAa	gtg	ıttgı	cgt	ago	cato	ctcc	agt	ttt	agt	tte	gtcg
		26:									_
NtA	DC2	:									
								. <b></b> .			т
GA.		c									
	٠g.	263	39								
	710	*	*	*	*	*	*	*	*		
*	*	* 1	+ +	**							
	720										
	DC1										
		gttt	++	,	+						
	tct	5000		<u></u>	.aat	<u>.a</u> cc		ayı	.cgg	luga	itgt
		678									
\T+- 7\	DC2										
					4	_					
		• • • •	• • •	<u>aat</u>	aat	<u>a</u>	• • •	• • •	• • •	• • •	• • •
• • •											
771	2	682									

Figure 7(h)

W. t	abecum				tabacun	aivshhsilifeavsass-hecssehlbegglosmaitlhedaladyanl	44
		mpalgccvdatvspplgyafsrdsslpapefffsgvpptmsaagsig	46	W.	sylvestris	Aivshesvlifeavsstt-trsqelssvdlqsfveklanddaradyrnl	44
	ylvastris	MPALGCCVDAAVSPPPGYSFLMDSSLPAPEIFPSGVPPSTNTAVATTTTT			esculentum	AIVSHHSVLITEAVSSTT-TRSQELSSVDLQSFVEKLNDDANADYRNL	44
	sculentum	MPALGCCVDAAVSPPPGYSI LADSSLPAFELI PROVPPSTNTAVATTTTT  MPALAC-VDTSFVPP-AYAFSDTAGDVF LPASSPTSA		۸.	theliens	AIVSHESVLIFEAVEADK-PHVHQATPGDIQFLLEG- NEEARANYEDL 434	
A. E	haliana 41	MENDAC-ADISTALL-VINGSDIMSDALINGSELSM		а.	max	AIVSHMSVLIFEAVGTSS-TN-GGGAPPALSAHYLAEELSIDYGYL	43
6.		NEVLACCYDAAAPPGYAFAGDISFPAPIAFT-GVPPATADD-THRENN			sative	AKASYHSKIILEALSAIPEPKDDEDEATTEOLHGRIRDLSSKLOPTGLSM	39
	etiva	NAXNYG			celi	AVTAINTYLVSNIIGVERNEYTVPTAPAEDAPRALCSHWETWOEN	40
1. 6		MSDBHSMGLPSEAGEHGVLASHQEVANS	28				
			,	N.	tabacum	BAAAVAGEYETCVLYSDQLKQRCVD-QFKEG\$LGIEHLAA	<b>∀D</b>
N. t	abecum					SICD 487	
					sylvestris	SAAAIRGEYDTCVLYADOLKORCVE-OFKDGDLDIEQLAAVDGICD	49
	ylvestris	hws panssaly sidgmgapy it twested is vaphet dtlphquidll kvv			esculentum	SAAAIRGEYDTCVLYADQLEGRCVE-QFEDGDLDIEQLAAVDGICD	49
	sculentum	Ewspanisalysidgmgapyftvmssgdisvkphgtdtlphqlidllkvv			theliane max	Yaavhrodhescllyvdolkorove—gykegylsieolasvd————glce Selafrodyetclvyteenkerove—ofrogtvoheolaave———-glce	47
A. t	haliens	RWSPSLSSSLYRIDGWGAPYFIANSSGNISVRPHGSETLE			sative	SSHAVKIRRGIK-MYKLGKK	40
		LETY 90			coli	HEFGTRASLAE-WINDSCHOLNDINIGYSSGIFSLOERAMAEQLYLSHCH	43
6		HUS PS LEAALYNVDGWGGFYF AVNTAGNISVAFHGSDTVSHQLIDLLKIV	96				
	etive	DVYHVEGWGEPYFAVNKDGHLCVRIYGRETLPGQEIDVLSVI	48				
î. :		SCHASICILATYNIAMWIRDIYYDVNELGHI SVCPDPDVPRARVDLAQLV		N.	tabarus	FVSKANGAADPIRTYNVNLSIFTSIPDFNAFGOLFPI	52
			)	N.	sylvestris	TVSKAIGASDFVRTYNVKLSIFTSVPDFWAIDOLFPI	52
			1	L-	esculentum	FVSKAIGASDPVRTTHVWLSIFTSVPDFWAIDQLFPI	52
N. 1	abacus	KKASDPKH8GGLGLQLPLVVRFPDVLHORLESLQSAFDLAVH9QGYGAHY	145 /	A.	theliene	WVLKAIGASDFVIT?	
N	ylvestris	KKASDPINLGGLGLQF PLVVRF PDI LKNRLESLQSVFDYAVQSQGYEAHY	150			YNINLSVYTSIPDLMGIDQLFPI 516	
L	sculentum	KKASDPKNLGGLGLQF PLVVRFPDI LICHKLESLQSVFDYAVQSQGYEAHY	150	٥.	MAX	Lvrkavgaaesvra	51
A. 1	haliene				setive	LSKSVTTDAHTIYNYPDNLSVFSLHPDYWGICHLFPH	45
		KKVTGPK880GLGLQLPLTVRFPDVLKHKLECLQSAFDYA	IKBQGY 1	E.	coli	eackordecompanys it depositementalianas aproched describe	50
		DSHY 140	***				
6. 1		KRASDPKSLGGLSLQLPLIARFPDVLKRLESLQSAFDYAIQSGGYESHY	146 97 1		*******	VPIHRLDERPAVRGILEDLTCDEDGKVDRFIGGESSLOLHELGENGD	
A. 1	METAN	EQATEADGTGK-KLQFPNILAFPDVLANRINSLHTAFANAIKYTQYGGVY			tabacum	A LTDRITTER LAANGE PROFICE BOOKANKE INCHESTORING	37
X. (	:011	KTREAQGORLPALICFPQILQHRLRSINAAFRARESYGYNGDY			sylvestris esculentum	VPIHKLDERPVVRGILSDLTCDSDGKIDKFIGGESSLPHKLGSNGG VPIHKLDERPVVRGILSDLTCDSDGKIDKFIGGESSLPHKLGSNGG	57
					theliene	VPINKLDORPGARGILBDLTCDSDGKINKFIGGIS	37
	abacum	OGVYPVRCHODRIVVEDIVRIGSSI RIGLEAGSKI ELLLANSCLCRGSAE	195	٠.	CHELIANA	STATHETHOUGS 263	
	rylvestris	OGVYPVKCNODRFVVEDIVKFGSGFRFGLEAGSKPELLLANSCLCKGSHE		۵.	mex	I PIHRLDEKPEVRGILEDLTCDSDGKIDKFINGESSLPLHIM	56
	sculentum	OGVY PVKCNODRFVVEDIVKFG8GFRFGLEAGSRPELLLANSCLCKGSHE			pativa	MPVSRLDERPTHRATLYDVTCDSDGKYDRFIRDTETHPLHPLDPRLG	30
	heliene				goli	LPLEGLDOVPERRAVLLDITCDSDGAIDHYIDGDGIATTHPHPEYDPENP	55
		QGVYPVKCHQDRFVVEDIVKFG88FRFGLEAG8KPEILLA	MSCLCK				
		GSPD 190					
6. 1	ax	QGVY PVKCHQDRFVVEDTVKFGS PFRFGLEAGSKPELLLANSCLCRGNPD			tabecus	GGGYYLDHF LGGAYEEALGGLINILFGG PSVVRVVQSDSAHEFAMS	61
A. 1	etive .	ogvi pvkvnonkovvodnyhi gydksygleagskpelliansclikakpg			mylvestris	GGGDGGK-YYLONG LGGAYELALGGLHNLFGGF5VLRV5QSD5FH5FAVT	62
E.	col1	flyypikynghrrvieslinggeplgleagskaelhavlahaghtr			esculentum	gggdggk-yylght lggayeealgglhnleggesvlavsgsdsphstavt	62
				A.	thelians	G	
				_		GR-YTLGHTLGGAYEEALGGVINLFGGPSVVRVSQSDGPHSFAVT 608	
	tabacum	GLLVCNGFKDAEYISLALVARKLHLNTVIVLEQEEELDLVIDISKKHAVR GLLVCNGFKDAEYISLALVARKLHLNTVIVLEQEEELDLVIDISKKHAVR			BAX Sativa	EGGRTYY LGNT LGGAY EEALGGVHN LFGG PSVVRVSQSDG PHSFAVT GYYVAV LLTGAY QEALSNION LFGG PSLVRVVGTGNGGAPNYE	60
	rylvestris	GILVENGE KDAEYISLALVARKLALMIVIVLEQESKLDLVIDISKANAVR			coli	HIGENACALÖET PAGGANTEGOLEVALAAG 1900-24EAE	54
	esculentum thaliane	APPACIACI EDUR I I BROTTA VENEZULA I A TROCER PRIME A I DI BENNEZA E	230	••	6011		39
^.	TOR 4 TORIS	AFLVCHGFEDAEYISLALLGAKLALNTVIVLEQEEKLDLV	TRLBOX				
		101VR 240		N.	tabacus	rsvpgpscadvlramchepelitetliodrazeflEquedkg	65
6. 1		GLLICHGFKDAEYISLALVANKLALHTVIVVEGEEEVDLIVELSKELCIK			wylvestris	CAVPGPSCADVLRAMOHEPELMYETLIGHRAEEFVHNDDEGEEDEG	66
	pativa	AYLVCHGYKDBAYVALALAARANGLNVIIVLEHEEELDIVIEESSKLGVE			esculentum	CAVPGPSCADVLAANGHEPELHTETLIGHAEEFVHNDDEQEEDKG	66
	coli	<b>SVIVCHGYKDREYIRLALIGEKHGHKVYLVIEKHSEIATVLDEAERLHVV</b>	216	λ.	theliens		
						Rav PGQ8 Sadvlranghe pe line of Lightae endit kogs:	LGT)
				_		EEED 658	
	tabecun	PVIGLRARLATKHSCHPOSTSGEKGKYGLTTTQIVRVVKKLEESCHLDCL			MAX	RAYPGPSCGDYLRYHQHQPELHTETLHHRAQEYVSHINAAA	64
	sylvestris	PVIGLRAKLRTKHBGHFG8T8GEKGKFGLTTTQIVRVVKKLEEBGHLDCL			sativa soli	aaligstteligtvsydvkodissvieerare Ledegdtvadmigyvoldpktlltgfrdgvkktdldaelooo	57
	esculentum	PVIGLRAKLRYKHIGHTGSTSGERGRYGLYTTQIVRVVKKLEESGMLDCL	300	••	6011	PROPERTY AND A STORE AND A VALUE OF THE PROPERTY OF THE PROPER	64
۸.	theliene	PVIGLBAKLRTRHSCHFGSTSGERGEFGLTTTQIVRVVRK	1.000000				
		LDCL 290		w.	tabanus	LAIASLASSLAGS-FHRHPYLVAPASCCFTAVTANNGGYNTYTEDEN	70
G. :		PVIGLRAKLRTRIBGHFGGIFRRAGKFGLTTARVLRVVENILDLAGGLDCL			sylvestris	LAFASLASSLAQS-FROMPYLVTNSSCCLTA-AANNGGY-YYCNDENIVG	71
	estive	PYIGVRAKLLTKI POHTGSTAGKHGKTGLPAKKIYEVAKKLKALNKLHML			esculentum	LAFASLASSLAGS-FROMFYLVTMSSCCLTA-RANNGGY-YYCNDENIVG	71
	poli	Prigvrarlasgeschiqebggersktglaatgvigivet laeagridsi			theliens	DEFINIVABILDAS-	_
						Propipy Late Qas Pensleaaienlg Pyycoedv 706	
					MAX	LINAGLART-FORMPYLLBLSSFVADOVARA	67
	tabecum	QLINTHIGSQIPSTALLADOVGEAAQIYCELIRLGAMMKFIDTGGGLG			estiva .	nkvwenverlvesglhthpyladykpppha	60
	sylvestris	QLIMTHIGSQIPSTALLADGVGEAAQIYCELVRLGASHKYIDCGGLG		ĸ.	coli	GLYGYTYLEDE	66
	esculentum	QLLHYHIGSQIPSTALLADGVGLAAQIYCELVALQAGNXYIDCGGGLG	348				
A.	theliene	QLINTHIGSQIPSTELLSDGVAEAAQLYCELVAL			tabacum	-AADSATGEDEIWSY-CTA	72
۵.		GAMMIVIDIGGES 338 OLIMPHIGSQIPSTALLADOVGEARQIYCILVRL—GAMMIVIDIGGES			sylvestris	VGAESAAAEKELMPY-CVA	73
		KLLHPHYGSHI PTTDIVFKAASEASDIYCALVKEYGVETHTTLDCGGGLG			esculentum	VGAESAAAEELWFY-CVA	73
	sativa coli	QLLHFHLGSQMANIRDIATGVRESARFYVELHRLGVNIQCFDVGGGLG			theliene	YDYI5-A	
						713	
				G.	max	VPAAQDLGEQWSY	69
N.	tabacum	IDYDGYRSCDSDVSVGYGIQEYASTVVQAVQYVCDRRGVRHPVICSESGR	393		mativa		
N.	tabacum sylvastris	IDYDGTKSCDSDCSVGYGLQEYASTVVQAVRFVCDRIGWXXHPVICSESGR	393 398		sativa coli		
N. L.	sylvestris esculentum		393				
N. L.	sylvestris	IDYDGTRSCDSDCSVGYGLQEYASTVVQAVALVCDARONXHFYICSISGA IDYDGTRSCDSDCSVGYGLQEYASTVVQAVALVCDARONXHFYICSISGA	393 398 390				
N. L.	sylvestris esculentum	IDYDGTKSCDSDCSVGYGLQEYASTVVQAVĀIVCDRKWVKHFVICSESGR IDYDGTKSCDSDCSVGYGLQEYASTVVQAVĀIVCDRKWVKHFVICSESGR IDYDGSKSGESDLSVAYSLKIYAEAVVASVRVVCDRSSVK	393 398 390				
N. L. A.	sylvestris esculentum thelians	IDTDOTESCHSUGGELDEVASTVVÖAVÄRVEDRUNVEHTVICSESGR IDTDOTESCHSUGGEVASTVVÖAVÄRVEDRUNVEHTVICSESGR IDTDOSESGESDLSVAYSLEITAEAVVASVRVVCDRSSVE ESGR 388	393 398 398 DEPVICS				
N. L. A.	sylvestris esculentum thelians max	IDYDOTESCOSICS WGYGLOF YASTYVÖAVÄY VCDROWNIN Y CSESGR IDYDGTHSCOSICS WGYGLOF YASTYVÖAVÄT VCDROWNIN Y CSESGR IDYDGSROGESOLS WASLETYARAWASVAVVAN Y CORDSSWI ESGR 388 IDYDGSRSCOSIDIS WETGLETYARAWYNIN Y COVERS-SVRHIV I CSESGR	393 391 393 393				
N. L. A.	sylvestris esculentum thelians	IDTDOTESCHSUGGELDEVASTVVÖAVÄRVEDRUNVEHTVICSESGR IDTDOTESCHSUGGEVASTVVÖAVÄRVEDRUNVEHTVICSESGR IDTDOSESGESDLSVAYSLEITAEAVVASVRVVCDRSSVE ESGR 388	393 398 398 DEPVICS				

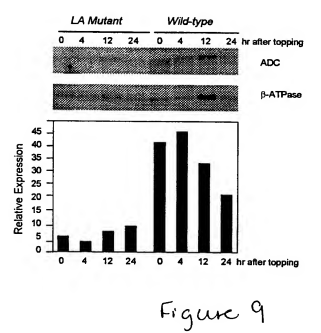


Figure 10

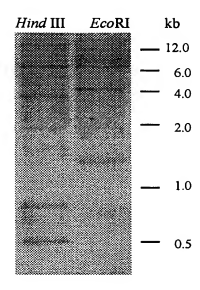


Figure 11

WO 00/67558

ODC2 ODC1			A AAGAAGAAAA TACTACGTAG A AAAAACAGTT TTTTTTTATT		
ODC2 ODC1			C AAAGGGAACA AGAGAAACAT G ATTCCTTTAA CAATGGCTTC		
ODC2 ODC1 pODC2 pODC1			T TEGATEGECEG GECAAACAAT T TEGATEGETEG GECAAACAAT H A G Q T I V	CATCGTTTCC GGGTTGAACC	
ODC2 ODC1 pODC2 pODC1		TCCTACAGCG GCGGCGGCG	G CGGAAAACGG CACCAGAAAA G CGGAAAACGA CACCAGAAAA A E N G T R K D		TGCCCTACAA GATTTCATGT
ODC2 ODC1 pODC2 pODC1	TATCAATCAT AACTCAAAAA		C TTTTTACGTG CTAGACTTGGCC TTTTTACGTG CTAGATTTGGPPFYVLDL		
ODC2 ODC1 pODC2 pODC1	CCCAAATATC CGTCCATTTT		T GAACCGTCGT TCCTTTCAAT T GAATCGTTGT TCCTTTCAAT P F P S F L S 1 S L	TTTATCTGCT ATGGGCTCAA	ATTTTGATTG TGCTAGCCGA
ODC2 ODC1 pODC2 pODC1		ATCTCTTGGC AATAAGAAA	G ACCGTATTGT TTTCGCAAAT A GAGAGAGGTC AATGGGTTAC D R I V F A N R E R S M G Y		GAAATTAATA TTGGGGTTGT
ODC2 ODC1 pODC2			T TTACAAGATC CGAAAGCATC A AATTTGGAGG AGTTTCTCTT V Y K I R K H		
ODC2 ODC1 pODC2	ATGAATAAAG CGAAGAACAC		C GCGCTTCCAG AAGAAGTCGAAACTCGAAACTCTTTTTCAACCAAAATT TCAATTTTTTCAACCAAAATT TCAATTTTTTCAACCAAACCAAAACCAAAACCAAAACCAAAACCAAAACCAAAA	TAAACGTTTT CTTTCTTGGT	
ODC2 ODC1 pODC2		ATTITTATTT TATTTATTA	T CAAACGCTTA TCTCGGCGCC A ATAGATTTAA CATAGTTTT S N A Y L G A	TTTACTCAAA ATAATATAT	TCATTTTTT ATTCGTCACT
ODC2 ODC1 pODC2			G GTTTACATCC GGCCACCAG G TTTGGCTGAT TGTTAAATAJ G F T S G H Q		
ODC2 ODC1 pODC2			A CCGGGTCGGT TTTTTGCAG	CGTATGTATT CAGCCAAAA1	
ODC2 ODC1 pODC2		ACCACGGGAC TCAGGGAAT	C TGTACGGTTC GATGAACTGT C CCTTACACCT TCTCCCCGGT L Y G S M N C		AGTTTGTTTT CGAAGACCAA
ODC2 ODC1 pODC2			C GAAAACGTTT CCGACGACTC A AAAAAGGTGA CTTGGAACAC S K T F P T T		
ODC2 ODC1 poDC2	TAATCCCTAT TTCAAATTTG	TCACTTTAAT TGGAAAAAC	TG GTTTTTCCTA ATATGGGTGG TT CTTTCACCCA CAATCCATA . V F P N M G J	A CAACACATTA TCTTTTGGAG	GTGTAAAAAG GTGATGTGAC
ODC2 ODC1 pODC2		GGGCTATTAA TAAGAATTC			1465 T TGATGGTTTT TTCCTTTTTT T TGGATATTAT TGTGTTTGGG
ODC2 ODC1				AATTCTCTTG TATGCC	A signal 1565 TGCAAGG ATTTGCTAAT CCCTCTGAGT CTTCTGATAG
ODC2 ODC1		AAGTTTTTAA AATTAGTTT			1665 T TGTGTGACTA TAAAAGCATC G AAGGATTTAA TCACATATGT

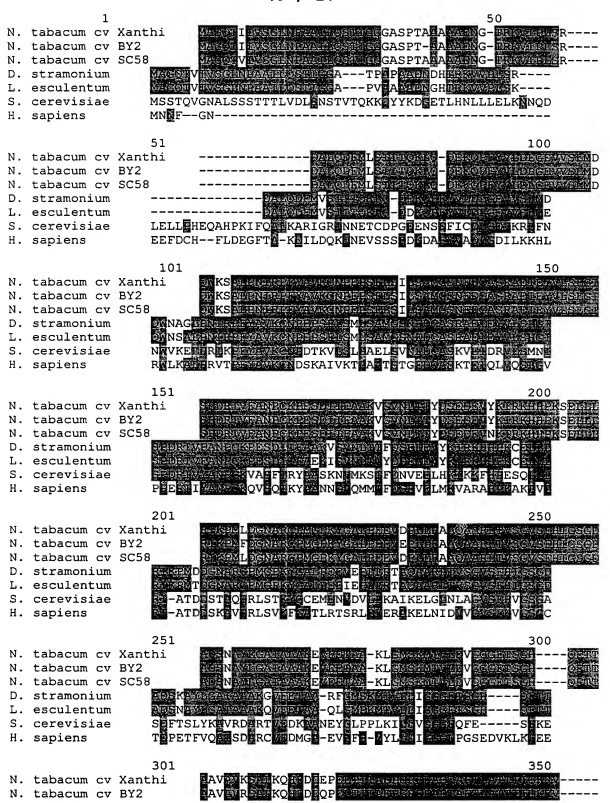
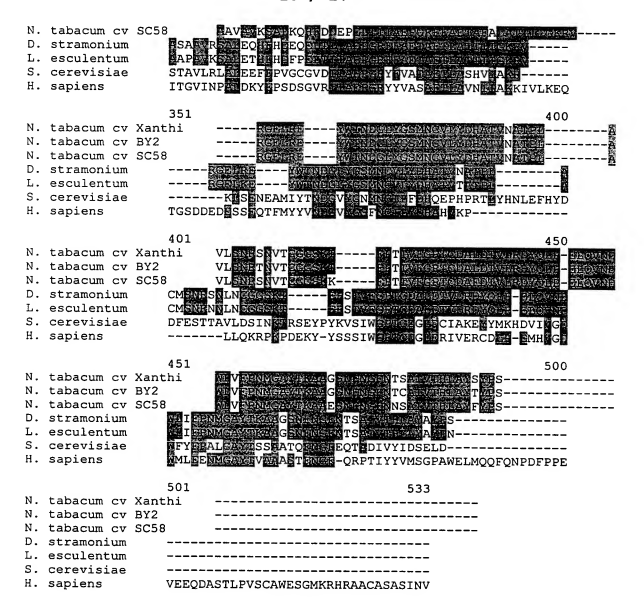
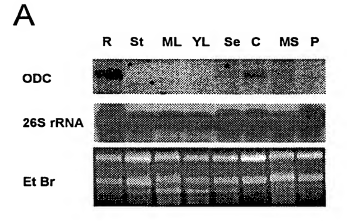


Figure 13 (a)





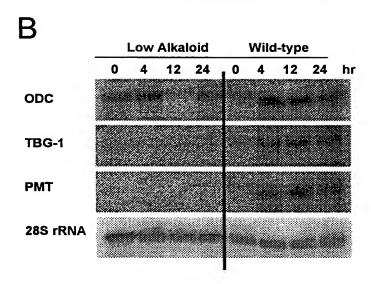


Figure 14

#### SEQUENCE LISTING

```
<110> Timko P, Michael
<120> Regulation of Gene Expression in Tabacco for
     Manipulation of Plant Growth and Secondary Metabolism
<130> 4981*239
<140>
<141>
<160> 26
<170> PatentIn Ver. 2.0
<210> 1
<211> 1120
<212> DNA
<213> Plant
<400> 1
ctgagttgac aagaacaatt cctggtgaat cagatggatg aagataatag aggtgggtgg 60
aatctataac caaagcagct ggttgagtga ctgtgcgagt tgcagaaaca attgaagggt 120
catttgtgga atttggggcc atttcaaagg aaaaagaaaa gatgacttag cattaataaa 180
tcaaattaaa ataaggctta gcgttaaaat caaaggaaat ggcaagcctg gctcctggag 240
caatqcttct qaqqacaqta qtaaaaacaa tatcagacaa aaagtaaagt tgtattattt 300
agcttgagga taaagtatgt cattagtttt gtgagagatt tggtgtcctc tacaatgatt 360
gttgaagtcc ctatttatag ctatacacag gaaacaaaat cctaggatca agcccctctt 420
aaatgacaat aatggggtta atgatgaata tgtagcggca tgacatgaat gccaaaattc 480
tccgcaacga ctatttattt aatattgagg aatatttttt attaaatact atctggtgac 540
aagcattegt ttgetteegt tgattaegtt gattttggga tetaetetat accaacegaa 600
gccgttgtcc ttgatcttcg ctttcattta attcatcttc cgtctgcctc cgatttcaca 660
agtcatgcac ccattcaatt atttaatgga aaccaatttt accctataca aatggtacat 720
cattegteaa ataetttaet tggatataaa caattttgee egaggagtaa acagatgega 780
agaaagaaag cagacgatta aagaaatttt taaaaaagga gagagaaatg aacacacaa 840
tgtactaata aaattagggt actactttac taataattgg acagagacta aattcatatt 900
ttagttccaa aatgtctcgg gcagtccaac catgcacgtt gtaatgattt tttaactcta 960
ttatatcgag ttgcgccctc cactcctcgg tgtccaaatt gtatataaat gcatatgtgt 1020
ctattgggag tgtacatcaa gctttcataa agtacaaatc gtaatacttg ttgaaacata 1080
                                                                   1120
atactttctc ttctccaatt tgtttagttt aattttgaaa
<210> 2
<211> 3091
<212> DNA
<213> Plant
<400> 2
```

ctgagttgac	aagaacaatt	cctggtgaat	cagatggatg	aagataatag	aggtgggtgg	60
aatctataac	caaagcagct	ggttgagtga	ctgtgcgagt	tgcagaaaca	attgaagggt	120
catttgtgga	atttggggcc	atttcaaagg	aaaaagaaaa	gatgacttag	cattaataaa	180
tcaaattaaa	ataaggctta	gcgttaaaat	caaaggaaat	ggcaagcctg	gctcctggag	240
caatgcttct	gaggacagta	gtaaaaacaa	tatcagacaa	aaagtaaagt	tgtattattt	300
agcttgagga	taaagtatgt	cattagtttt	gtgagagatt	tggtgtcctc	tacaatgatt	360
gttgaagtcc	ctatttatag	ctatacacag	gaaacaaaat	cctaggatca	agcccctctt	420
aaatgacaat	aatggggtta	atgatgaata	tgtagcggca	tgacatgaat	gccaaaattc	480
tccgcaacga	ctatttattt	aatattgagg	aatattttt	attaaatact	atctggtgac	540
aagcattcgt	ttgcttccgt	tgattacgtt	gattttggga	tctactctat	accaaccgaa	600
gccgttgtcc	ttgatcttcg	ctttcattta	attcatcttc	cgtctgcctc	cgatttcaca	660
agtcatgcac	ccattcaatt	atttaatgga	aaccaatttt	accctataca	aatggtacat	720
cattcgtcaa	${\tt atactttact}$	tggatataaa	caattttgcc	cgaggagtaa	acagatgcga	780
agaaagaaag	cagacgatta	aagaaatttt	taaaaaagga	gagagaaatg	aacacacaca	840
-					aattcatatt	
-					tttaactcta	
					gcatatgtgt	
					ttgaaacata	
					tatctaccaa	
					accataatgg	
					acgggacaat	
					ctggttggtt	
					atgcatattt	
					taattaaaaa	
					ctctacacat	
					gagaagttac	
					aatatttaa	
_					gcatgtgtac	
					acacacagag	
					catcccaaac	
					aatgcttcgt	
					tgatgtaagt	
					ttttctaaaa	
_					atttttccct	
					tggtgcgtat	
					taccttttgg	
					tttgtaaagg	
					attggtactc	
					cggtaattga	
					cattctttga	
					aaagcatttg	
					ttaagggctc	
					tctctctc	
					tcagttccaa	
					ttttcaactg	
					tctactgaag	
					gctcaagtca ctcacaattt	
agtccaaatt	agcacctctc	aayttetaca	accergacge	aactccatat	CCCacaaccc	2000

ctttttcct attgtacttt atgttcttcg tcaaatttta taattaactc ttttcaaatt 2940 gtctttttt ttttcagatt cacaaagcag cattcatttt gccatctttc gccagaagta 3000 tgatcgagtc ttaatcaact gattaatgaa tactggtggt acaatcattg gaccaagatc 3060 aataagtgaa agacgtattg tatgagaatt c 3091

<210> 3

<211> 353

<212> PRT

<213> Plant

<400> 3

Met Glu Val Ile Ser Thr Asn Thr Asn Gly Ser Thr Ile Phe Lys Ser 1 5 10 15

Gly Ala Ile Pro Met Asn Gly His His Asn Gly Thr Ser Lys His Gln
20 25 30

Asn Gly His Lys Asn Gly Thr Ser Glu Gln Gln Asn Gly Thr Ile Ser 35 40 45

Leu Asp Asn Gly Asn Glu Leu Leu Gly Asn Ser Asn Cys Ile Lys Pro 50 55 60

Gly Trp Phe Ser Glu Phe Ser Ala Leu Trp Pro Gly Glu Ala Phe Ser 65 70 75 80

Leu Lys Val Glu Lys Leu Leu Phe Gln Gly Lys Ser Asp Tyr Gln Asp 85 90 95

Val Met Leu Phe Glu Ser Ala Thr Tyr Gly Lys Val Leu Thr Leu Asp 100 105 110

Gly Ala Ile Gln His Thr Glu Asn Gly Gly Phe Pro Tyr Thr Glu Met 115 120 125

Ile Val His Leu Pro Leu Gly Ser Ile Pro Asn Pro Lys Lys Val Leu 130 135 140

Ile Ile Gly Gly Gly Ile Gly Phe Thr Leu Phe Glu Met Leu Arg Tyr 145 150 155 160

Pro Thr Ile Glu Lys Ile Asp Ile Val Glu Ile Asp Asp Val Val Val 165 170 175

Asp Val Ser Arg Lys Phe Phe Pro Tyr Leu Ala Ala Asn Phe Asn Asp 180 185 190

Pro Arg Val Thr Leu Val Leu Gly Asp Gly Ala Ala Phe Val Lys Ala

3

195 200 205

Ala Gln Ala Glu Tyr Tyr Asp Ala Ile Ile Val Asp Ser Ser Asp Pro 210 215 220

Ile Gly Pro Ala Lys Asp Leu Phe Glu Arg Pro Phe Phe Glu Ala Val 225 230 235 240

Ala Lys Ala Leu Arg Pro Gly Gly Val Val Cys Thr Gln Ala Glu Ser 245 250 255

Ile Trp Leu His Met His Ile Ile Lys Gln Ile Ile Ala Asn Cys Arg 260 265 270

Gln Val Phe Lys Gly Ser Val Asn Tyr Ala Trp Thr Thr Val Pro Thr 275 280 285

Tyr Pro Thr Gly Val Ile Gly Tyr Met Leu Cys Ser Thr Glu Gly Pro 290 295 300

Glu Ile Asp Phe Lys Asn Pro Val Asn Pro Ile Asp Lys Glu Thr Ala 305 310 315 320

Gln Val Lys Ser Lys Leu Ala Pro Leu Lys Phe Tyr Asn Ser Asp Ile 325 330 335

His Lys Ala Ala Phe Ile Leu Pro Ser Phe Ala Arg Ser Met Ile Glu 340 345 350

Ser

<210> 4

<211> 711

<212> DNA

<213> Plant

<400> 4

gaattcaatg gagaaggaa atattccag tgtaaacaca agtgaatgaa gagaagccaa 60 aataatctct atcattcaag ccttaggtgg agattaaaaa aattattac tttcttatca 120 aagtaatagg tgatcaacag ctttcgtaaa acgtcattag gagaatatta taatctctt 180 tatgctgaag aacccacata aggaagatca taaaatacat gactttcaga tgacttcttg 240 gagctttatt tttaaagagt ggctagctgg tcagcaaga ggtgctcgtc agatatcata 300 aaattttact attattgtt ttaagaggga gatggggcac acatgcttgt gacaaaagta 360 agaggaagaa aggagacaga agaggaagaa atgaggacaca acatatacta acaaaattt 480 actaataatt gcaccgagac aaacttatat tttagttcca aaatgtcagt ctaaccctgc 540

acgttgtaat gaattttaa ctattatat atatcgagtt gcgccctcca ctcctcggtg 600 tccaaattgt atttaaatgc atagatgtt attgggagtg tacagcaagc tttcggaaaa 660 tacaaaccat aatacttct cttcttcaat ttgtttagtt taattttgaa a 711

<210> 5 <211> 3129 <212> DNA

<213> Plant

<400> 5

gaattcaatg gagaaggaaa atatttccag tgtaaacaca agtgaatgaa gagaagccaa 60 aataatetet ateatteaag eettaggtgg agattaaaaa aattatttae tttettatea 120 aagtaatagg tgatcaacag ctttcgtaaa acgtcattag gagaatatta taatctcttt 180 tatgctgaag aacccacata aggaagatca taaaatacat gactttcaga tgacttcttg 240 gagetttatt tttaaagagt ggetagetgg teageaaaga ggtgetegte agatateata 300 aaattttact attatttgtt ttaagaggga gatggggcac acatgcttgt gacaaaagta 360 tcaaagaaaa tttttaaaat ggagagagaa atgagcacac acatatacta acaaaatttt 480 actaataatt gcaccgagac aaacttatat tttagttcca aaatgtcagt ctaaccctgc 540 acgttgtaat gaatttttaa ctattatatt atatcgagtt gcgccctcca ctcctcggtg 600 tccaaattgt atttaaatgc atagatgttt attgggagtg tacagcaagc tttcggaaaa 660 tacaaaccat aatactttct cttcttcaat ttgtttagtt taattttgaa aatggaagtc 720 atatetacea acacaaatgg etetaceate tteaagaatg gtgeeattee catgaacgge 780 caccaaaatg gcacttctga acacctcaac ggctaccaga atggcacttc caaacaccaa 840 aacgggcacc agaatggcac tttcgaacat cggaacggcc accagaatgg gacatccgaa 900 caacagaacg ggacaatcag ccatgacaat ggcaacgagc tactgggaag ctccgactct 960 attaagcctg gctggttttc agagtttagc gcattatggc caggttagta ctaagaaagc 1020 aactcaaatq catcqqcctc ttqttqctac taaatataqa qaqctatcat acttttaggg 1080 actaactaaa aaggaaagat tatcacaggg acgaagtgag cagttaactt cgcatattat 1140 cagacgcatt aatttgaaat aatcgaattt tgcaggtgaa gcattctcac ttaaggttga 1200 gaagttacta ttccagggga agtctgatta ccaagatgtc atgctctttg aggtaattaa 1260 tattctaata cacatgcttt aatttaaagt gatactttta atttactttt agtttattgc 1320 atgtgcacgt acagtcagca acttatggga aggttctgac tttggatgga gcaattcaac 1380 atacagagaa tggtggattt ccatacactg aaatgattgt tcatctacca cttggttcca 1440 tcccaaaccc aaaaaaggtt ttgatcatcg gcggaggaat tggttttaca ttattcgaaa 1500 tgcttcgtta tccttcaatc gaaaaaattg acattgttga gatcgatgac gtggtagttg 1560 atqtaaqtca aacttctttt acccacataa aqaaaatgat ttagattgca attcttttta 1620 caggtatcca gaaaattttt cccttatctg gcagctaatt ttaacgatcc tcgtgtaacc 1740 ctagttctcg gagatggtgc gtatatgata gtctcgtttt atattttatt tcacttgatt 1800 tttacctttt tttqtqqtta attaatcatc taccattggt tctctttacc ttcaggagct 1860 gcatttgtaa aggctgcaca agcgggatat tatgatgcta ttatagtgga ctcttctgat 1920 cccattggta cgctattact atttaatacc aagactattc ttattaaata agctactaag 1980 aaactaattg aataattaat aaacgtaact gtaattgatt tctaaaataa tatatataat 2040 ttcaggtcca gcaaaagatt tgtttgagag gccattcttt gaggcagtag ccaaagccct 2100 taggccagga ggagttgtat gcacacaggc tgaaagcatt tggcttcata tgcatattat 2160 taagcaaatc attgctaact gtcgtcaagt ctttaagggt tctgtcaact atgcttggac 2220 aaccgttcca acatatccca cgtattcttt ttctctctct ctcttcctgt ctttttcgat 2280

gcaatgtaaa tttataaaat tggaagtccg ttttactttt ctatagacgt agatcctaaa 2340 attgtcaaga aatggagaat tgacttacaa gaaaaatcaa cttcttttca tttactattc 2400 tttttggtga caaactttac ttattatttc gttctaaaat gaaaatttat ttttatattt 2460 taaaataatt tagctttaaa cttttaattt tacttgttat atttttaata aaaaagattt 2520 atagtcaaat aaatgttgtg accatataaa aacctccgca tttttaagat cataagtttc 2580 agagtcaaac gagttaattt atttttagta tgccggtgcg gagtcaaatt atgtcataaa 2640 aattgaaacg gagtgagaac atttttattt cgagtaaact ttcaaggtat tgtgtttaat 2700 ttcaagtgat actgatcaat gatgtcttaa atattttgat ttcagcggtg tgatcggtta 2760 tatgctctgc tctactgaag ggccagaagt tgacttcaag aatccagtaa atccaattga 2820 caaagagaca actcaagtca agtccaaatt aggacctctc aagttctaca actctgatgt 2880 aacttcatat ctcacaattt ctttttccgt tttactgtat gttcttcgtc aaattttata 2940 actaactctt ttcatattgt ctttttttc agattcacaa agcagcattc attttaccat 3000 ctttcgccag aagtatgatc gagtcttaat caagtgaata atgaacactg gtagtacaat 3060 cattggacca agatcgagtc ttaatcaagt gaataaataa gtgaaatgcg acgtattgta 3120 3129 ggagaattc

<210> 6

<211> 375

<212> PRT

<213> Plant

<400> 6

Met Glu Val Ile Ser Thr Asn Thr Asn Gly Ser Thr Ile Phe Lys Asn 1 5 10 15

Gly Ala Ile Pro Met Asn Gly His Gln Asn Gly Thr Ser Glu His Leu 20 25 30

Asn Gly Tyr Gln Asn Gly Thr Ser Lys His Gln Asn Gly His Gln Asn 35 40 45

Gly Thr Phe Glu His Arg Asn Gly His Gln Asn Gly Thr Ser Glu Gln
50 55 60

Gln Asn Gly Thr Ile Ser His Asp Asn Gly Asn Glu Leu Leu Gly Ser
65 70 75 80

Ser Asp Ser Ile Lys Pro Gly Trp Phe Ser Glu Phe Ser Ala Leu Trp 85 90 95

Pro Gly Glu Ala Phe Ser Leu Lys Val Glu Lys Leu Leu Phe Gln Gly 100 105 110

Lys Ser Asp Tyr Gln Asp Val Met Leu Phe Glu Ser Ala Thr Tyr Gly 115 120 125

Lys Val Leu Thr Leu Asp Gly Ala Ile Gln His Thr Glu Asn Gly Gly 130 135 140

6

Phe 145	Pro	Tyr	Thr	Glu	Met 150	Ile	Val	His	Leu	Pro 155	Leu	Gly	Ser	Ile	Pro 160
Asn	Pro	Lys	Lys	Val 165	Leu	Ile	Ile	Gly	Gly 170	Gly	Ile	Gly	Phe	Thr 175	Leu
Phe	Glu	Met	Leu 180	Arg	Tyr	Pro	Ser	Ile 185	Glu	Lys	Ile	Asp	Ile 190	Val	Glu
Ile	Asp	Asp 195	Val	Val	Val	Asp	Val 200	Ser	Arg	Lys	Phe	Phe 205	Pro	Tyr	Leu
Ala	Ala 210	Asn	Phe	Asn	Asp	Pro 215	Arg	Val	Thr	Leu	Val 220	Leu	Gly	Asp	Gly
Ala 225	Ala	Phe	Val	Lys	Ala 230	Ala	Gln	Ala	Gly	Tyr 235	Tyr	Asp	Ala	Ile	Ile 240
Val	Asp	Ser	Ser	Asp 245	Pro	Ile	Gly	Pro	Ala 250	Lys	Asp	Leu	Phe	Glu 255	Arg
Pro	Phe	Phe	Glu 260	Ala	Val	Ala	Lys	Ala 265	Leu	Arg	Pro	Gly	Gly 270	Val	Val
Cys	Thr	Gln 275	Ala	Glu	Ser	Ile	Trp 280	Leu	His	Met	His	Ile 285	Ile	Lys	Gln
Ile	Ile 290		Asn	Cys	Arg	Gln 295		Phe	Lys	Gly	Ser 300	Val	Asn	Tyr	Ala
Trp 305	Thr	Thr	Val	Pro	Thr 310	Tyr	Pro	Thr	Gly	Val 315	Ile	Gly	Tyr	Met	Leu 320
Cys	Ser	Thr	Glu	Gly 325		Glu	Val	Asp	Phe 330		Asn	Pro	Val	Asn 335	Pro
Ile	Asp	Lys	Glu 340		Thr	Gln	Val	Lys 345		Lys	Leu	Gly	Pro 350	Leu	Lys
Phe	Tyr	Asn 355		Asp	Ile	His	Lys 360	Ala	Ala	Phe	Ile	Leu 365		Ser	Phe
Ala	Arg 370	Ser	Met	. Ile	e Glu	Ser 375									

<210> 7

<211> 1134 <212> DNA <213> Plant

#### <400> 7

gctgtacaaa aggatgtctc aaatcatttg gaatattaat tctgcaatca acaagaaata 60 ccccactatt aagacccatt atcactggca caaaaattat gagatcatta aacatcttaa 120 acctgtccct atttggaaga gtgtggtatg ggagatgcct cccagggagt acctaaagct 180 gaatactgat ggaagtttta acaaacaaat tgggaaagca gggattggag ggattctcag 240 agatgaagag ggaggetttg teatggettt ttegatgeet ataatetata ataacateag 300 tgaagcagaa ttgaaagcca tcaagtatgg gtgtgaatgg tgcaaataca aaggaatatc 360 aaacttcatt gtggaaactg actcgaggat gatctatgac atactacaga ccaaaaatct 420 aagcaacaac aagttgaaac aagagaccga gaaattaatg gagattctgg acacctgcag 480 gacacctgtt acccattgcc ttcgcgaagc aaatcaagtg gcagactggt ttgctaaaga 540 ggccaccaga gctaacgaag gtatcactca tacagatttt agacaggtat caaaagcggc 600 caaqqqccct ttcttcatqq atatqtqqca qqtcccttat tttagaatta gatatgaaaa 660 atctaatttt tttttgtaag ttaattctgt gtatagtgag aggaaatcgt ctaatatgta 720 tttttgccca tagactcttc ctctccttag gtaaaaaggt agctccgagg taaggtttat 780 gttcccctca gtgtaacctt tttttgttta tataatagac atggtatggg tccagctaaa 840 ccccaacac cacaggggat agatacctgg gtgattggtt tatttttaa aaaaaaaaac 900 tttactaata attqcacqqa qacaaaactt atattttagt tccaaaatga cagtccaacc 960 atgcacgttg taatgatttt ttaactctat tatatcgagt tccgccctcc actcctcggt 1020 gtccaaattg tatttaaatg catagatatg tttattggga gtgtacatca agctttcaga 1080 aaatacaaac cataatactt tctcttctcc aatttgctta gtttaatttg gaaa 1134

<210> 8 <211> 3269 <212> DNA <213> Plant

<400> 8

gctgtacaaa aggatgtctc aaatcatttg gaatattaat tctgcaatca acaagaaata 60 cccactatt aagacccatt atcactggca caaaaattat gagatcatta aacatcttaa 120 acctgtccct atttggaaga gtgtggtatg ggagatgcct cccagggagt acctaaagct 180 qaatactgat ggaagtttta acaaacaaat tgggaaagca gggattggag ggattctcag 240 agatgaagag ggaggctttg tcatggcttt ttcgatgcct ataatctata ataacatcag 300 tgaagcagaa ttgaaagcca tcaagtatgg gtgtgaatgg tgcaaataca aaggaatatc 360 aaacttcatt gtggaaactg actcgaggat gatctatgac atactacaga ccaaaaatct 420 aagcaacaac aagttgaaac aagagaccga gaaattaatg gagattctgg acacctgcag 480 gacacctgtt acccattgcc ttcgcgaagc aaatcaagtg gcagactggt ttgctaaaga 540 ggccaccaga gctaacgaag gtatcactca tacagatttt agacaggtat caaaagcggc 600 caagggccct ttcttcatgg atatgtggca ggtcccttat tttagaatta gatatgaaaa 660 atctaatttt tttttgtaag ttaattctgt gtatagtgag aggaaatcgt ctaatatgta 720 tttttgccca tagactcttc ctctccttag gtaaaaaggt agctccgagg taaggtttat 780 gttcccctca gtgtaacctt tttttgttta tataatagac atggtatggg tccagctaaa 840 ccccaacac cacaggggat agatacctgg gtgattggtt tatttttaa aaaaaaaaac 900 tttactaata attgcacgga gacaaaactt atattttagt tccaaaatga cagtccaacc 960 atgcacgttg taatgatttt ttaactctat tatatcgagt tccgccctcc actcctcggt 1020

```
qtccaaattq tatttaaatq cataqatatq tttattggga gtgtacatca agctttcaga 1080
aaatacaaac cataatactt tctcttctcc aatttgctta gtttaatttg gaaaatggaa 1140
qtcatatcta ccaacacaaa tggctctact atcttcaaga atggtgccat tcccatgaac 1200
ggttaccaga atggcacttc caaacaccaa aacggccacc agaatggcac ttccgaacat 1260
cggaacggcc accagaatgg gatttccgaa caccaaaacg gccaccagaa tggcacttcc 1320
gagcatcaga acggccatca gaatgggaca atcagccatg acaacggcaa cgagctacag 1380
ctactgggaa gctccaactc tattaagcct ggttggtttt cagagtttag cgcattatgg 1440
ccaggttagt actaagaaag aaactcaaat gcatcgtact cttgtattct gctttgcgta 1500
taatttagat gatggtgttt gactaagcac tgagtttaaa aataaaaagt ttaaagttaa 1560
attgttacta tagagagcta tatctttagg aactaactaa aaaggaaaaa ttatcacata 1620
aaattgggat gaagtaagca gttaacttcg catattattc gacacattaa tttgaaataa 1680
atcgaatttt gcaggtgaag cattctcact taaggttgag aagttactat tccaggggaa 1740
gtctgattac caagatgtca tgctctttga ggtaattaat taatactaat agtcaagctc 1800
atgtatgatt atatttaaag tggtattttt cgtttatttt taatttattg cacgtgtacg 1860
tacagtcagc aacatatggg aaggttctga ctttggatgg agcaattcaa cacacagaga 1920
atggtggatt tocatacact gaaatgattg ttcatcttcc acttggttcc atcccaaacc 1980
ctaaaaaggt tttgatcatc ggcggaggaa ttggttttac attattcgaa atgcttcgtt 2040
atcctacaat cqaaaaaatt qacattqttq aqatcqatqa cgtggtagtt gatgtaagtc 2100
aaacttcttt tactcacata aaaaaatgat ttagattctt atttttctaa aagaattaaa 2160
acaaaatttt ccgttttaca ggtatctaga aaatttttcc cttatcttgc tgctaatttt 2220
agggatecte gtgtaaceet agteettgga gatggtgegt atttgataat etegttttta 2280
ttttatcttt tacttttatt ttatttaatt tttacctttt tgtgtgtggt taattcacct 2340
gccattggtt ctttttattt caggggctgc atttgtaaag gccgcacaag caggatatta 2400
tgatgctatt atagtggact cttctgatcc cattggtact ctattactac ttaataccaa 2460
qactattett attaaataaq etaetaataa aegtaaetet gatagtttte taaaataata 2520
taatttcagg tccagcaaaa gacttgtttg agaggccatt ctttgaggca gtagccaaag 2580
ccctaaggcc aggaggagtt gtatgcacac aggctgaaag catttggctt catatgcata 2640
ttattaagca aatcattgct aactgtcgtc aagtctttaa gggctctgtc aactatgctt 2700
ggactactgt tccaacatat ccaacgtatt tttctctctc tcttcctata aaattggaag 2760
ttttqattct ataattqtca aqaaatqqaq aatcagttcc aagaaaaacc aaattctttt 2820
cttttactct tcaaggtgtg tttaagtttt ttaaactgat actgatcaat tattttgatt 2880
tcagcqqtqt gattqqttat atgctctqtt ctactqaagg accagaagtt gacttcaaga 2940
atccagtaaa tccaattgac aaagagacaa ctcaagtcaa gtccaaatta gcacctctca 3000
agttctacaa ctctgatgta acttcatatc tcaatttctt ttttcttatt gtactttatg 3060
ttcttagtca aattttataa ttaactcttt tcaaattgtc tttttttttc agattcacaa 3120
agcagcattc attttgccat ctttcgccag aagtatgatc gagtcttaat caagtgacta 3180
atgaatactg gcggtacaat cattggacca agatcgagtc ttaatcaagt gaataaataa 3240
                                                                  3269
gtgaaatgcg acgtattgta taagaattc
```

```
<211> 381
<212> PRT
<213> Plant
<400> 9
Met Glu Val Ile Ser Thr Asn Thr Asn Gly Ser Thr Ile Phe Lys Asn
1 5 10 15
```

<210> 9

9

Gly Ala Ile Pro Met Asn Gly Tyr Gln Asn Gly Thr Ser Lys His Gln Asn Gly His Gln Asn Gly Thr Ser Glu His Arg Asn Gly His Gln Asn Gly Ile Ser Glu His Gln Asn Gly His Gln Asn Gly Thr Ser Glu His Gln Asn Gly His Gln Asn Gly Thr Ile Ser His Asp Asn Gly Asn Glu Leu Gln Leu Leu Gly Ser Ser Asn Ser Ile Lys Pro Gly Trp Phe Ser Glu Phe Ser Ala Leu Trp Pro Gly Glu Ala Phe Ser Leu Lys Val Glu Lys Leu Leu Phe Gln Gly Lys Ser Asp Tyr Gln Asp Val Met Leu Phe Glu Ser Ala Thr Tyr Gly Lys Val Leu Thr Leu Asp Gly Ala Ile Gln His Thr Glu Asn Gly Gly Phe Pro Tyr Thr Glu Met Ile Val His Leu Pro Leu Gly Ser Ile Pro Asn Pro Lys Lys Val Leu Ile Ile Gly Gly Gly Ile Gly Phe Thr Leu Phe Glu Met Leu Arg Tyr Pro Thr Ile Glu Lys Ile Asp Ile Val Glu Ile Asp Asp Val Val Asp Val Ser Arg Lys Phe Phe Pro Tyr Leu Ala Ala Asn Phe Ser Asp Pro Arg Val Thr Leu Val Leu Gly Asp Gly Ala Ala Phe Val Lys Ala Ala Gln Ala Gly Tyr Tyr Asp Ala Ile Ile Val Asp Ser Ser Asp Pro Ile Gly Pro Ala Lys Asp Leu Phe Glu Arg Pro Phe Phe Glu Ala Val Ala Lys Ala Leu

Arg Pro Gly Gly Val Val Cys Thr Gln Ala Glu Ser Ile Trp Leu His 280 275 Met His Ile Ile Lys Gln Ile Ile Ala Asn Cys Arg Gln Val Phe Lys 295 Gly Ser Val Asn Tyr Ala Trp Thr Thr Val Pro Thr Tyr Pro Thr Gly 310 315 320 Val Ile Gly Tyr Met Leu Cys Ser Thr Glu Gly Pro Glu Val Asp Phe 325 330 335 Lys Asn Pro Val Asn Pro Ile Asp Lys Glu Thr Thr Gln Val Lys Ser 340 345 350 Lys Leu Ala Pro Leu Lys Phe Tyr Asn Ser Asp Ile His Lys Ala Ala 355 360 Phe Ile Leu Pro Ser Phe Ala Arg Ser Met Ile Glu Ser 370 375 <210> 10 <211> 469 <212> DNA <213> Plant <400> 10 gtcgacctct gattccacaa gtcatgcacc cattcaatta tttaatggaa accaatttta 60 ccctgtacaa atggtacaaa tactttcctt ggataaaaac aattttgcct aaggagtaaa 120 cagatgcgaa gtaagaaagc agacgactaa agaaaatttt aaaaaaggag agagaaatga 180 gcacacaca gtactaataa aattagggta ctactttact aataattgga cagagactaa 240 attcatattt tagttccaaa atgtctcggg cagtccaacc atgcacgttg taatgagttt 300 ttaactctat tatctcgagt tgcgccctcc actcctctgt gtccaagttg tatataaatg 360 catatatgtc tattgggagt gtacagcgag ctttcataaa gtacaaatca taatacttgt 420 <210> 11 <211> 3001 <212> DNA <213> Plant <400> 11 gtcgacctct gattccacaa gtcatgcacc cattcaatta tttaatggaa accaatttta 60 ccctgtacaa atggtacaaa tactttcctt ggataaaaac aattttgcct aaggagtaaa 120 cagatgcgaa gtaagaaagc agacgactaa agaaaatttt aaaaaaggag agagaaatga 180 gcacacaca gtactaataa aattagggta ctactttact aataattgga cagagactaa 240 attcatattt tagttccaaa atgtctcggg cagtccaacc atgcacgttg taatgagttt 300

ttaactctat	tatctcgagt	tacaccctcc	actcctctqt	gtccaagttg	tatataaatg	360
					taatacttgt	
					tggaagtcat	
-					tgaatggcca	
					aacaccaaaa	
					tttccgaaca	
					atgggatttc	
					accagaatgg	
					tactgggaaa	
-					caggttagta	
					ataagaagtt	
					caaaaaatga	
					ttgcaggtga	
agcattctcc	cttaaggttg	agaagttact	atttcagggg	aagtctgact	accaagatgt	1140
catgctcttt	gaggtaaata	atattctaat	acacatgctt	taatatgaat	aaatactttt	1200
					aaggttttga	
	_				gaaatgattg	
					ggcggaggaa	
					gacattgttg	
					aaaaaatgat	
ttagattgct	tctttttatt	tttctaaaag	aataaatata	ttctctctta	gttttaaaca	1560
aaattctctt	tcttacaggt	atctagaaaa	tctttccctt	atctcgcagc	taattttaat	1620
gatcctcgtg	taaccctcgt	tctcggagat	ggtgcgtatt	tataatctcg	tttttgtttt	1680
atcttttatt	tttatttcat	ttaatttacc	tttttgtgtg	tggttaattt	acccgtcatt	1740
ggttctcttt	catttcaggg	gctgcatttg	taaaggctgc	acaagcagga	tattatgatg	1800
ctattatagt	ggactcttct	gatcccattg	gtactctatt	actacttaat	accaagacta	1860
atcttattga	ataagctact	aataaactgt	aattgatttc	taaaataata	taatttcagg	1920
tccagcaaaa	gatttgtttg	agaggccatt	ctttgaggca	gtagccaaag	ccctaaggcc	1980
aggaggagtt	gtatgcacac	aggccgaaag	catttggctt	catatgcata	ttattaagca	2040
aatcattgct	aactgtcgtc	aagtctttaa	gggctctgtc	aactacgctt	ggactactgt	2100
tccaacatat	cccacgtatt	ttctctctct	ctctcttcat	ctttgaaaat	tgaaaatcct	2160
gactactttc	cttcctttga	ttcctcggtt	aaaggggcgt	agatcataag	attttcaaga	2220
aatagataat	gacgtccaag	aaaaactaac	ttcttttcat	ttactattct	ttttggtgac	2280
aaactttatt	tattatttcg	ttctaaagag	aaaatttatt	tttatattt	aaaataattt	2340
					gtcaaataaa	
					ccaaatgagt	
					gtaatggagt	
					ttcaactgat	
					tatgctctgc	
					caaagagaca	
					aacttcatat	
					aattaactct	
					tctttcgcca	
					tcattggacc	
aagatcgagt	cttaatcaag	tgaataaata	agtgaaatgc	cgacgtattg	tatgagaatt	
С						3001

<211> 419

<212> PRT

<213> Plant

<400> 12

Met Glu Val Ile Ser Thr Asn Thr Asn Gly Ser Thr Ile Phe Lys Asn 1 5 10 15

Gly Ala Ile Pro Met Asn Gly His Gln Ser Gly Thr Ser Lys His Leu 20 25 30

Asn Gly Tyr Gln Asn Gly Thr Ser Lys His Gln Asn Gly His His Asn 35 40 45

Gly Thr Ser Glu His Arg Asn Gly His Gln Asn Gly Ile Ser Glu His 50 55 60

Gln Asn Gly His Gln Asn Gly Thr Ser Glu His Arg Asn Gly His Gln 65 70 75 80

Asn Gly Ile Ser Glu His Gln Asn Gly His Gln Asn Gly Thr Ser Glu 85 90 95

His Gln Asn Gly His Gln Asn Gly Thr Ser Glu Gln Gln Asn Gly Thr
100 105 110

Ile Ser His Asp Asn Gly Asn Glu Leu Leu Gly Asn Ser Asn Ser Ile 115 120 125

Lys Leu Gly Trp Phe Ser Glu Phe Ser Ala Leu Trp Pro Gly Glu Ala 130 135 140

Phe Ser Leu Lys Val Glu Lys Leu Leu Phe Gln Gly Lys Ser Asp Tyr 145 150 155 160

Gln Asp Val Met Leu Phe Glu Ser Ala Thr Tyr Gly Lys Val Leu Thr 165 170 175

Leu Asp Gly Ala Ile Gln His Thr Glu Asn Gly Gly Phe Pro Tyr Thr 180 185 190

Glu Met Ile Val His Leu Pro Leu Gly Ser Ile Pro Asn Pro Lys Lys 195 200 205

Val Leu Ile Ile Gly Gly Gly Ile Gly Phe Thr Leu Phe Glu Met Leu 210 215 220

Arg Tyr Pro Thr Ile Glu Lys Ile Asp Ile Val Glu Ile Asp Asp Val

225 230 235 240

Val Val Asp Val Ser Arg Lys Ser Phe Pro Tyr Leu Ala Ala Asn Phe 245 250 255

Asn Asp Pro Arg Val Thr Leu Val Leu Gly Asp Gly Ala Ala Phe Val 260 265 270

Lys Ala Ala Gln Ala Gly Tyr Tyr Asp Ala Ile Ile Val Asp Ser Ser 275 280 285

Asp Pro Ile Gly Pro Ala Lys Asp Leu Phe Glu Arg Pro Phe Phe Glu 290 295 300

Ala Val Ala Lys Ala Leu Arg Pro Gly Gly Val Val Cys Thr Gln Ala 305 310 315 320

Glu Ser Ile Trp Leu His Met His Ile Ile Lys Gln Ile Ile Ala Asn 325 330 335

Cys Arg Gln Val Phe Lys Gly Ser Val Asn Tyr Ala Trp Thr Thr Val 340 345 350

Pro Thr Tyr Pro Thr Gly Val Ile Gly Tyr Met Leu Cys Ser Thr Glu 355 360 365

Gly Pro Glu Val Asp Phe Lys Asn Pro Ile Asn Pro Ile Asp Lys Glu 370 375 380

Thr Thr Gln Val Lys Ser Lys Leu Ala Pro Leu Lys Phe Tyr Asn Ser 385 390 395 400

Asp Ile His Lys Ala Ala Phe Ile Leu Pro Ser Phe Ala Arg Ser Met 405 410 415

Ile Glu Ser

<210> 13

<211> 1636

<212> DNA

<213> Plant

<400> 13

ggcacgagat cagatccaat tetetetgt getteeette tetgetetea aattetteag 60 atetacaaag tetteeteat tetegaggg cagacatgga aacteteteg tetaceteag 120 agteagteaa tgaaggecae eeegacaage tetgegacea ggteteggat geaattettg 180

```
atgcttgctt agaacaggat ccagaaagca aggttgcatg tgaaacctgc acaaagacaa 240
acatggttat ggtctttgga gagatcacaa ccaaggccac tgttgactat gagaagatag 300
tgcqtqacac atqcaqagqc attgggttca cctcagcaga tgttggcctt gacgctgaca 360
actgcaaggt tottgtcaac atcgagcagc agagccctga cattgcccaa ggtgttcacg 420
gtcatcttac caagaaacca gaagagattg gagctggtga ccaaggtcac atgtttggct 480
atgccactga tgaaacccca gagctcatgc cccttaccca tgtttgggcc actaagcttg 540
gtgccaagct taccgaagtg aggaagaaca agacttgccc atggctcaga ccagatggca 600
agacccaagt tactgttgag tacaagaacg acaatggtgc catggtccca attagagttc 660
acactgttct catctcaact caacatgacg aaactgtcac aaacgaccag attgcccagg 720
acttgaaaga gcatgtgatc aaacctgtga tcccatctca gtaccttgat gagaatacca 780
tettecacet caacecatea ggtegetteg teateggtgg accaeagga gatgetggae 840
ttaccggcag gaaaattatc attgacacct acggaggctg gggtgcccat ggaggaggtg 900
ctttctcagg aaaggaccct actaaggtgg acaggagtgg tgcttatatt gttagacagg 960
cagcaaagag tgtggtcgcc tcaggacttg ctcgccgctg tattgtgcag gtttcttatg 1020
ctatcggtgt ggctgaacca ctttccgtgt ttgttgacac ttacaagact ggaacaattc 1080
cagacaagga tattttgact ctgatcaagg agaactttga cttcaggcct ggaatgatgt 1140
caatcaacct tgacttgtta agaggaggca acttcaggta ccagaagact gcagcttatg 1200
qtcactttqq ccqtqatqac cccqacttct catgggagac tgtcaaggtc ctcaagccaa 1260
aagcttaagt gaggtgtagc cttttggcca ttatttttct tgcagaccaa taaacaagct 1320
tcatcatatc atgcattggt ggcaggagaa gagaatttgt gtctccattg gaggattcta 1380
tgagctctga gtcattgaac attgttattt ttctttcttt ttttttcacc cttttctgca 1440
gtaccttatt tttattttgt tactgttaag tagcagtgat ttaagttttc cctgttaagt 1500
agcagtgatt taagttttcc ctgttaagta gctggaatta agtttccatg ttctatcata 1560
ttatatgtga acttgtcaat tatctcctga ggtgaaagag tccttcaggg aatagtttaa 1620
                                                                  1636
aaaaaaaaa aaaaaa
```

<210> 14

<211> 390

<212> PRT

<213> Plant

<400> 14

Met Glu Thr Phe Leu Phe Thr Ser Glu Ser Val Asn Glu Gly His Pro 1 5 10 15

Asp Lys Leu Cys Asp Gln Val Ser Asp Ala Ile Leu Asp Ala Cys Leu 20 25 30

Glu Gln Asp Pro Glu Ser Lys Val Ala Cys Glu Thr Cys Thr Lys Thr 35 40 45

Asn Met Val Met Val Phe Gly Glu Ile Thr Thr Lys Ala Thr Val Asp
50 55 60

Tyr Glu Lys Ile Val Arg Asp Thr Cys Arg Gly Ile Gly Phe Thr Ser 65 70 75 80

Ala Asp Val Gly Leu Asp Ala Asp Asn Cys Lys Val Leu Val Asn Ile

85	90	95

PCT/US00/12450

WO 00/67558

Glu Gln Gln Ser Pro Asp Ile Ala Gln Gly Val His Gly His Leu Thr Lys Lys Pro Glu Glu Ile Gly Ala Gly Asp Gln Gly His Met Phe Gly Tyr Ala Thr Asp Glu Thr Pro Glu Leu Met Pro Leu Thr His Val Trp Ala Thr Lys Leu Gly Ala Lys Leu Thr Glu Val Arg Lys Asn Lys Thr Cys Pro Trp Leu Arg Pro Asp Gly Lys Thr Gln Val Thr Val Glu Tyr Lys Asn Asp Asn Gly Ala Met Val Pro Ile Arg Val His Thr Val Leu Ile Ser Thr Gln His Asp Glu Thr Val Thr Asn Asp Gln Ile Ala Gln Asp Leu Lys Glu His Val Ile Lys Pro Val Ile Pro Ser Gln Tyr Leu Asp Glu Asn Thr Ile Phe His Leu Asn Pro Ser Gly Arg Phe Val Ile Gly Gly Pro His Gly Asp Ala Gly Leu Thr Gly Arg Lys Ile Ile Ile Asp Thr Tyr Gly Gly Trp Gly Ala His Gly Gly Gly Ala Phe Ser Gly Lys Asp Pro Thr Lys Val Asp Arg Ser Gly Ala Tyr Ile Val Arg Gln Ala Ala Lys Ser Val Val Ala Ser Gly Leu Ala Arg Arg Cys Ile Val Gln Val Ser Tyr Ala Ile Gly Val Ala Glu Pro Leu Ser Val Phe Val Asp Thr Tyr Lys Thr Gly Thr Ile Pro Asp Lys Asp Ile Leu Thr Leu 

Ile Lys Glu Asn Phe Asp Phe Arg Pro Gly Met Met Ser Ile Asn Leu

340 345 350

Asp Leu Leu Arg Gly Gly Asn Phe Arg Tyr Gln Lys Thr Ala Ala Tyr 355 360 365

Gly His Phe Gly Arg Asp Asp Pro Asp Phe Ser Trp Glu Thr Val Lys 370 375 380

Val Leu Lys Pro Lys Ala 385 390

<210> 15

<211> 1596

<212> DNA

<213> Plant

### <400> 15

ggcacgaggg gaacaagaga aacatcatat tattgaatcc ctagtttctt ttctttccct 60 ttgattcctt cctctcattt acctctctt tttcttcctt tgtttggatg gccggccaaa 120 caatcatcgt ttccgggttg aacccggcgg ccattcttca gtccacaatt ggcggcggag 180 cttctcctac agcggcggcg gcggcggaaa acggcaccag aaaagtcatc cctctctcaa 240 gagatgcctt acaagatttc atgttatcaa tcataaccca aaaattacaa gatgagaaac 300 aaccttttta cgtgctagac ttgggtgagg ttgtttctct tatggaccaa tggaaatctg 360 ctctcccaaa tatccgtcca ttttacgctg ttaaatgtaa ccctgaaccg tcgttccttt 420 caattttatc tgctatgggc tcaaattttg attgtgctag ccgagctgaa attgagtatg 480 ttttatctct tggcatttca cctgaccgta ttgttttcgc aaatccatgc aaaccggaat 540 ccgatattat ttttgcagca aaagttgggg tgaatcttac aacctatgat tctgaagacg 600 aggtttacaa gatccgaaag catcacccga aatccgaact cttgctccgc atcaagccca 660 tgctcgacgg caacgcgaga tgcccaatgg gcccgaaata cggcgcgctt ccagaagaag 720 tegacceget geteegggea geteaageeg ecegteteae egtateegge gteteattee 780 acateggtag eggagatgee gatteaaaeg ettatetegg egecatagee geggetaagg 840 aagtgtttga aacagctgct aaactcggga tgtcgaaaat gactgttcta gacgtcggcg 900 gegggtttac atceggecac cagtteacaa eegeegeegt egeegttaaa teagetttaa 960 aacaacatt cgatgacgaa ccggagttga caatcatagc tgaaccgggt cggttttttg 1020 cagagacggc gtttactttg gcaacgacga ttatagggaa aagagtgagg ggtgaattga 1080 gggagtattg gattaacgac gggctgtacg gttcgatgaa ctgtgtactt tacgaccatg 1140 cgacggtgaa tgcaacgccg ttagctgttc tgtcgaatcg tagtaacgtt acctgcggcg 1200 ggtcgaaaac gtttccgacg actgtgtttg ggcccacttg tgatgctctt gatactgttt 1260 taagggatta ccagttaccg gagctgcagg ttaatgattg gctggttttt cctaatatgg 1320 gtgcttatac taaagctgct gggtccaatt ttaatggatt taatacttcc gccattgtta 1380 ctcacctcgc ttattcttat ccaagctgat gaaccacctg tattaggaat tactaccgtg 1440 gttttgatgg ttttttcctt ttttgggtat ctttttttta attttgttgt ttttggtagt 1500 aatttatatt ccaaatcagc ttgtaattct cttgtatgcc ataagaatgc aaggatttgc 1560 1596 taattgtgat tttctctaaa aaaaaaaaa aaaaaa

<210> 16

<211> 433

<212> PRT <213> Plant

<400> 16

Met Ala Gly Gln Thr Ile Ile Val Ser Gly Leu Asn Pro Ala Ala Ile 1 5 10 15

Leu Gln Ser Thr Ile Gly Gly Gly Ala Ser Pro Thr Ala Ala Ala Ala 20 25 30

Ala Glu Asn Gly Thr Arg Lys Val Ile Pro Leu Ser Arg Asp Ala Leu 35 40 45

Gln Asp Phe Met Leu Ser Ile Ile Thr Gln Lys Leu Gln Asp Glu Lys
50 55 60

Gln Pro Phe Tyr Val Leu Asp Leu Gly Glu Val Val Ser Leu Met Asp 65 70 75 80

Gln Trp Lys Ser Ala Leu Pro Asn Ile Arg Pro Phe Tyr Ala Val Lys 85 90 95

Cys Asn Pro Glu Pro Ser Phe Leu Ser Ile Leu Ser Ala Met Gly Ser 100 105 110

Asn Phe Asp Cys Ala Ser Arg Ala Glu Ile Glu Tyr Val Leu Ser Leu 115 120 125

Gly Ile Ser Pro Asp Arg Ile Val Phe Ala Asn Pro Cys Lys Pro Glu 130 135 140

Ser Asp Ile Ile Phe Ala Ala Lys Val Gly Val Asn Leu Thr Thr Tyr 145 150 155 160

Asp Ser Glu Asp Glu Val Tyr Lys Ile Arg Lys His His Pro Lys Ser 165 170 175

Glu Leu Leu Arg Ile Lys Pro Met Leu Asp Gly Asn Ala Arg Cys 180 185 190

Pro Met Gly Pro Lys Tyr Gly Ala Leu Pro Glu Glu Val Asp Pro Leu 195 200 205

Leu Arg Ala Ala Gln Ala Ala Arg Leu Thr Val Ser Gly Val Ser Phe 210 215 220

His Ile Gly Ser Gly Asp Ala Asp Ser Asn Ala Tyr Leu Gly Ala Ile 225 230 235 240

Ala Ala Lys Glu Val Phe Glu Thr Ala Ala Lys Leu Gly Met Ser

245 250 Lys Met Thr Val Leu Asp Val Gly Gly Phe Thr Ser Gly His Gln 260 265 270 Phe Thr Thr Ala Ala Val Ala Val Lys Ser Ala Leu Lys Gln His Phe 275 280 285 Asp Asp Glu Pro Glu Leu Thr Ile Ile Ala Glu Pro Gly Arg Phe Phe 295 300 Ala Glu Thr Ala Phe Thr Leu Ala Thr Thr Ile Ile Gly Lys Arg Val 305 310 315 Arg Gly Glu Leu Arg Glu Tyr Trp Ile Asn Asp Gly Leu Tyr Gly Ser 325 330 Met Asn Cys Val Leu Tyr Asp His Ala Thr Val Asn Ala Thr Pro Leu 345 Ala Val Leu Ser Asn Arg Ser Asn Val Thr Cys Gly Gly Ser Lys Thr 355 360 365

Phe Pro Thr Thr Val Phe Gly Pro Thr Cys Asp Ala Leu Asp Thr Val 370 375 380

Leu Arg Asp Tyr Gln Leu Pro Glu Leu Gln Val Asn Asp Trp Leu Val 385 390 395 400

Phe Pro Asn Met Gly Ala Tyr Thr Lys Ala Ala Gly Ser Asn Phe Asn 405 410 415

Gly Phe Asn Thr Ser Ala Ile Val Thr His Leu Ala Tyr Ser Tyr Pro 420 425 430

Ser

<210> 17

<211> 2074

<212> DNA

<213> Plant

<400> 17

tggtaactgg accgacgcga catttgtcgt atatgtctta atcgggctag tcgctgacaa 60

```
catcatccac caagtcaaag ttcggaaatt catatcgttt ctcatcatct tctatccgag 120
aaatgagggg actatctgta tacggtcaaa accgagtctg cccttcatat gactaatcga 180
gattagaaca taatggtcta aggttcatca ttataataac gagccatgat atagagttag 240
gttgtcaagc tcaagcccca gagagcgatc aatatcgaga tcgagccaag gttaaactcg 300
agacctagag accgatcaat accgagaccg accaagtcaa ggtcgagctc gagacccaga 360
gaccggtcaa gattgagatc ggccaagatc gagatcgagc caagaaatta aaaagtcgtt 420
atagccgcat ttagggagag aatctctgcg gaaatcacga cttgaatcag ggaaaaacta 480
attaattaat ctatcatgtg atccccacta tgtattttta attatactca aaatgggatt 540
ccccactat attaagagtg gttatcattt gtaatggaga gacatacaca cattcattct 600
gacatataca gaaaatagag caaatactat ccttttttgg cttttgatat ttagtcatat 660
tgtttcttct acccattgtt cttcactcaa tttggaggtg ataaaacttg aaggtttaag 720
ctaactagtc cattcgggtt gcattcattt cttttacaat aatttcgtca tcatttattt 780
attttctcaa ttgtactaag ttataccacg tatttttaga actgcgtata aattcaactc 840
tatccatttt tcgggtaaac accgaataca tagcacaata gcaccctcaa ttgcaaaagt 900
ccaaagccaa gggttcattc ctttctgaag aaatgagata gagaattgaa aatctaattt 960
agttatctaa atctttataa tttagccttc catataagaa aaaggaaaca aattaactga 1020
agaacaatag cctcgcatag atttaccttc tccatataaa ttttgtttat actcaatttt 1080
tttgcaaatg tgtctaaaat gataggactt gcaaattttt atttaacatt tcctactcct 1140
ctttaatttt caagaaatta attttaagca ttctcgattt gctctgcccg ctccgtcccg 1200
ttgccatctc tgactcggat aggacctcga ttgcaaaaat ccaaacccaa ggaaccttcc 1260
atacattaca taagccacaa aatagtaact attaaaaact accaatatat cctcaaatac 1320
tegegattat tteataceta acaegtttae ettatettet egtaatgaeg etaeattagt 1380
tagtgatata aaataccgaa tttaccacgc ggcaaccctc cgctgtctat ccacggcccg 1440
agagaatete ttageecec aaatacgaaa attaaettet agaattttat tttetggtta 1500
ttaccatgaa aataaagaaa aagagaaaag tcaagaaatt taattgggct aatactgggg 1560
tccactgccc agccacgcat ttccctccta tataaagcgt cgtcacctct catgcaaatc 1620
tcgctcactt cacagttgtt agtttcacgt tctcttctca attcccataa aagaaaccct 1680
teegttaggt tteegteeta ttttetette ttetaegett cetettetga tateaatate 1740
tgtatggtgt ttttcttgtt cgaattttag atttgttttg cctttaatac ctgtaacctt 1800
ataattctct gtttaaacca aaaacttagc ttcttctgaa gtcagggtgg ggatatttgg 1860
atcgtgtaag agtgtgttag aaggtgatta tcttttgatt cagttccttt tttgcttctt 1920
ttgagggggt agccggggcc tcggcctcgg cgggttttaa tagcccccat ctattacaac 1980
cattgggcaa aaacatcatt aaatctgtac aaagcaaacc cttaatttag tttaattttc 2040
                                                                  2074
tgtattcttt gattctttaa cagaagaaga agag
```

<210> 18

<211> 4321

<212> DNA

<213> Plant

### <400> 18

tggtaactgg accgacgga catttgtcgt atatgtctta ategggctag tcgctgacaa 60 catcatcac caagtcaaag ttcggaaatt catatcgtt ctcatcatct tctatccgag 120 aaatgagggg actatctgta tacggtcaaa accgagtctg cccttcatat gactaatcga 180 gattagaaca taatggtcta aggttcatca ttataataac gagccatgat atagagttag 240 gttgtcaagc tcaagccca gagagcgatc aatatcgaga tcgagccaag gttaaactcg 300 agacctagaa gattgagatc ggccaagatc gagatcgacc caagaatta aaaagtcgtt 420

atagccgcat	ttagggagag	aatctctgcg	gaaatcacga	cttgaatcag	ggaaaaacta	480
					aaatgggatt	
					cattcattct	
					ttagtcatat	
					aaggtttaag	
					tcatttattt	
					aattcaactc	
					ttgcaaaagt	
					aatctaattt	
					aattaactga	
					actcaatttt	
					tcctactcct	
					ctccgtcccg	
ttgccatctc	tgactcggat	aggacctcga	ttgcaaaaat	ccaaacccaa	ggaaccttcc	1260
atacattaca	taagccacaa	aatagtaact	attaaaaact	accaatatat	cctcaaatac	1320
tcgcgattat	ttcataccta	acacgtttac	cttatcttct	cgtaatgacg	ctacattagt	1380
tagtgatata	aaataccgaa	tttaccacgc	ggcaaccctc	cgctgtctat	ccacggcccg	1440
agagaatctc	ttagcccccc	aaatacgaaa	attaacttct	agaattttat	tttctggtta	1500
ttaccatgaa	aataaagaaa	aagagaaaag	tcaagaaatt	taattgggct	aatactgggg	1560
tccactgccc	agccacgcat	ttccctccta	tataaagcgt	cgtcacctct	catgcaaatc	1620
tcgctcactt	cacagttgtt	agtttcacgt	tctcttctca	attcccataa	aagaaaccct	1680
tccgttaggt	ttccgtccta	ttttctcttc	ttctacgctt	cctcttctga	tatcaatatc	1740
tgtatggtgt	ttttcttgtt	cgaattttag	atttgttttg	cctttaatac	ctgtaacctt	1800
					ggatatttgg	
					tttgcttctt	
					ctattacaac	
cattgggcaa	aaacatcatt	aaatctgtac	aaagcaaacc	cttaatttag	tttaattttc	2040
					gttgcgtaga	
					ttcccgcgcc	
					ttgggtctcc	
					tctccgttaa	
					accaggaaat	
					ggctcgggct	
					aatctctgca	
					aaggtgttta	
					tcgggtcgtc	
attccggttc	gggttggaag	ctgggtctaa	acccgagctc	ctgttagcca	tgagctgtct	2640
ctgcaggggc	agtgctgagg	gccttctcgt	ttgcaatggt	ttcaaggacg	ctgagtacat	2700
ttcgcttgct	ttggttgcaa	gaaagctcat	gttaaacact	gtaattgttc	ttgaacaaga	2760
					ccgtaattgg	
					ctggagaaaa	
aggtaagttt	gggcttacaa	cgacccaaat	tgttcgtgta	gtgaagaagc	tggaagaatc	2940
					tcccttcaac	
					taatccgtct	
tggtgcgggt	atgaagttca	ttgatactgg	aggtgggctc	ggaattgatt	atgatggtac	3120
					cctccacagt	
tgtccaggcg	gttcaatatg	tttgcgaccg	taagggcgtg	aagcacccag	tgatttgcag	3240
cgaaagtggc	agggcaattg	tttctcatca	ctcaattctg	attttcgaag	ccgtgtctgc	3300

ttctaqtcac tcatqttctt cttcacatct gtcttctggt ggcctccaat ccatggcgga 3360 gacgeteaat gaagatgeee ttgetgatta eegeaattta tetgetgetg eagttegtgg 3420 agagtacgag acgtgtgtac tttactctga tcagttgaaa cagagatgtg tggatcagtt 3480 taaagaaggg teettgggta tigaacatet tgetgetgtt gatageatet gtgattttgt 3540 atcaaaggct atgggggctg ctgatcctat ccgcacttac catgtgaatc tgtcaatttt 3600 cacttcaatt cctgattttt gggcctttgg tcaattgttt ccgattgttc caatacaccg 3660 tttaqatqaa aaqcctqcag taaggggaat attatcggac ttgacttgtg acagtgatgg 3720 gaaggttgat aagttcattg gtggcgaatc aagcttgcag ctgcatgaat tgggaagtaa 3780 tggcgatggt ggtgggtatt atctggggat gtttttgggt ggggcttatg aggaggcgct 3840 cggaggactc cacaacctgt ttggtggacc aagcgtggtg cgcgtggtgc agagcgatag 3900 cgctcacage ttegecatgt etegeteegt ceetggeeeg teetgegegg aegtgeteeg 3960 agcgatgcag cacgagcccg agctcatgtt cgagactctc aagcaccgtg cggaggaatt 4020 cttggaacaa gaagaagaca aagggctggc cattgcatct ttggccagca gcttagctca 4080 qtccttccat aacatgcctt accttgtggc gcctgcatct tgctgcttca ctgcagttac 4140 tgctaacaac ggtggctata actactatta cagtgatgag aatgcagcag attctgctac 4200 aggggaggat gagatttggt cctattgcac tgcttgaagt gttgtcgtag catctccagt 4260 tttagtttgt cgtcgaagtt gtctgttttt gaataatacc cttagttggt gatgttttc 4320

<210> 19

<211> 720

<212> PRT

<213> Plant

<400> 19

Met Pro Ala Leu Gly Cys Cys Val Asp Ala Thr Val Ser Pro Pro Leu
1 5 10 15

Gly Tyr Ala Phe Ser Arg Asp Ser Ser Leu Pro Ala Pro Glu Phe Phe 20 25 30

Thr Ser Gly Val Pro Pro Thr Asn Ser Ala Ala Gly Ser Ile Gly Ser 35 40 45

Pro Asp Leu Ser Ser Ala Leu Tyr Gly Val Asp Gly Trp Gly Ala Pro 50 55 60

Tyr Phe Ser Val Asn Ser Asn Gly Asp Ile Ser Val Arg Pro His Gly 65 70 75 80

Thr Asp Thr Leu Pro His Gln Glu Ile Asp Leu Leu Lys Val Val Lys
85 90 95

Lys Ala Ser Asp Pro Lys Asn Ser Gly Gly Leu Gly Leu Gln Leu Pro 100 105 110

Leu Val Val Arg Phe Pro Asp Val Leu Lys Asn Arg Leu Glu Ser Leu 115 120 125

Gln	Ser 130	Ala	Phe	Asp	Leu	Ala 135	Val	His	Ser	Gln	Gly 140	Tyr	Gly	Ala	His
Туг 145	Gln	Gly	Val	Tyr	Pro 150	Val	Lys	Cys	Asn	Gln 155	Asp	Arg	Phe	Val	Val 160
Glu	Asp	Ile	Val	Lys 165	Phe	Gly	Ser	Ser	Phe 170	Arg	Phe	Gly	Leu	Glu 175	Ala
Gly	Ser	Lys	Pro 180	Glu	Leu	Leu	Leu	Ala 185	Met	Ser	Cys	Leu	Cys 190	Arg	Gly
Ser	Ala	Glu 195	Gly	Leu	Leu	Val	Cys 200	Asn	Gly	Phe	Lys	Asp 205	Ala	Glu	Tyr
Ile	Ser 210	Leu	Ala	Leu	Val	Ala 215	Arg	Lys	Leu	Met	Leu 220	Asn	Thr	Val	Ile
Val 225	Leu	Glu	Gln	Glu	Glu 230	Glu	Leu	Asp	Leu	Val 235	Ile	Asp	Ile	Ser	Arg 240
Lys	Met	Ala	Val	Arg 245	Pro	Val	Ile	Gly	Leu 250	Arg	Ala	Lys	Leu	Arg 255	Thr
Lys	His	Ser	Gly 260	His	Phe	Gly	Ser	Thr 265	Ser	Gly	Glu	Lys	Gly 270	Lys	Phe
Gly	Leu	Thr 275	Thr	Thr	Gln	Ile	Val 280	Arg	Val	Val	Lys	Lys 285	Leu	Glu	Glu
Ser	Gly 290		Leu	Asp	Cys	Leu 295		Leu	Leu	His	Phe 300	His	Ile	Gly	Ser
Gln 305	Ile	Pro	Ser	Thr	Ala 310	Leu	Leu	Ala	Asp	Gly 315	Val	Gly	Glu	Ala	Ala 320
Gln	Ile	Tyr	Cys	Glu 325		Ile	Arg	Leu	Gly 330		Gly	Met	Lys	Phe 335	Ile
Asp	Thr	Gly	Gly 340		Leu	Gly	Ile	Asp 345		Asp	Gly	Thr	Lys 350	Ser	Cys
Asp	Ser	Asp 355		Ser	Val	Gly	Туг 360		Ile	Gln	Glu	Туг 365		Ser	Thr
Val	Val		Ala	Val	Gln	Tyr 375		Cys	Asp	Arg	Lys 380		Val	Lys	His

9ro 385	Val	lle	Cys	Ser	390	ser	GIÀ	Arg	Ala	395	vai	ser	nis	urs	400
Ile	Leu	Ile	Phe	Glu 405	Ala	Val	Ser	Ala	Ser 410	Ser	His	Ser	Cys	Ser 415	Ser
Ser	His	Leu	Ser 420	Ser	Gly	Gly	Leu	Gln 425	Ser	Met	Ala	Glu	Thr 430	Leu	Asn
Glu	Asp	Ala 435	Leu	Ala	Asp	Tyr	Arg 440	Asn	Leu	Ser	Ala	Ala 445	Ala	Val	Arg
Gly	Glu 450	Tyr	Glu	Thr	Cys	Val 455	Leu	Tyr	Ser	Asp	Gln 460	Leu	Lys	Gln	Arg
Cys 465	Val	Asp	Gln	Phe	Lys 470	Glu	Gly	Ser	Leu	Gly 475	Ile	Glu	His	Leu	Ala 480
				485					490				Gly	495	
_			500					505					Thr 510		
	-	515					520					525	Pro		
	530					535					540		Asp		
545					550					555			Glu		560
				565					570				Gly	575	
			580					585					Gly 590		
		595					600					605	Gln		
	610					615					620		Pro		
Ala 625		Val	Leu	Arg	Ala 630		Gln	His	Glu	Pro 635		Leu	Met	Phe	Glu 640

Thr Leu Lys His Arg Ala Glu Glu Phe Leu Glu Glu Glu Glu Asp Lys 645 650 655

Gly Leu Ala Ile Ala Ser Leu Ala Ser Ser Leu Ala Gln Ser Phe His 660 665 670

Asn Met Pro Tyr Leu Val Ala Pro Ala Ser Cys Cys Phe Thr Ala Val 675 680 685

Thr Ala Asn Asn Gly Gly Tyr Asn Tyr Tyr Tyr Ser Asp Glu Asn Ala 690 695 700

Ala Asp Ser Ala Thr Gly Glu Asp Glu Ile Trp Ser Tyr Cys Thr Ala 705 710 715 720

<210> 20

<211> 2118

<212> DNA

<213> Plant

### <400> 20

gaattootta tooggattto tggtacgcag actgtaatat ggagtoatot totootogat 60 tcgggattaa aattaggtga cttgggacac cctaaatctc ccaagtggcg actctgaaat 120 aaataaacaa atcccgtttc gattgtcctt aaattggaaa aaactccctt gtaccctccc 180 gggtacggaa aaaggaggtg tacagcaatg acccaaaact tttattgcta tacattttga 240 ggaatcaact tgatcaaaat ttatgggtga aattcaatgt ggtatgattt atattaggtc 300 ggactttagc agatgtggtc acttcaattt gcggcaaaaa taatgtacag ggataataat 360 aaaaagtact agaaatttga gtcataaagc tttttcaatt ttacaaaaga tattaagata 420 cttattaaat caaatgtact ttattaatgt aatagcatga aaaaacagcc tcatccgcct 480 gtcctcaccc cacaaaagg agatagagaa aggaaactaa tcttatttaa ttttccacat 540 tttaatcatc cgtataagaa agaagctaat taactgactt acaaactgaa tagatagcac 660 aatagcactc tcaattacaa aaatccaaag ccgagggtca ttcctttcat caagaaatta 720 gatagggaat ggaaaatata atttaattat ctgaatcttt ataatttatc cttccatata 780 agaaaaagga aacaaattaa ctgaagagca tatagcctcg catagattta ccttctccat 840 atgggtgggg aaaccgacaa accgcaccaa ttcgataatt cgagtcaaac tgaggaaaaa 900 aaaattcgac tatggtttgg tttgatttgg tttattgttg ggataaaaaa tcgatcataa 960 ttggtttggt ttggttttaa ctaaagaaag tcaaaccgaa accaaacaaa cccgacatta 1020 catatataaa ttttttagat atatttaata tataaatata cttgttgtga tgtaatttat 1080 aaatatttct taaaaatatt cataatttta tettttaaga tattattteg taettagaae 1140 ttttgaatgt ttcttactcc tctttaattt gttagaaatt aatttgaagg agtttgaatt 1200 tgctccgccc ccgctccgtc ccgttgccat ccctgactca ataggataac agcaatctcg 1260 attgcaaaaa tccaaaccca aggaaccttc ccaacattac ataagctaca aagtagagta 1320

gtttattaaa taactacaa tatacctca aattetegeg attattteat acctaacaeg 1380 cttaccttat cttetegtaa tgacgetaca ttagttggtg atataaaata cegaatttge 1440 cacgeggcaa teeteegetg tetateeaeg geeegagaga atetettage eeeceaaaga 1500 tgaaaattaa ettetagaat tttatttet ggttattace atgaaaataa ttaaataaaa 1560 aaaaagagaa aagtaaagat atttaattgg getaaaaetg gggteeaegg eeeaggeeaeg 1620 catteeete etaataaag egtegteaee teteatgeaa ateteegeta etaacaegtt 1680 gttagttea egteetete teaatteeea taacagaaae eetteegtea ggttteegte 1740 etattttee teatettee egttteetet tetgaaatea atatetgtat ggtgtttte 1800 ttgttegaat tagettette tgaagteagg gtggggatt ttggategt taagagtgtg 1920 ttagagggtg attatettt gatteagte eetattata aeettaaat eetetgttaa 1860 ageeteggee teggeggtt ttaatagee eeatetata eaettatag gggtageegg 1980 ggeeteggee teggeggtt ttaatagee eeatetata eaetattg geaaaaaca 2040 cattaaate gtacaaaaa aaceettaat ttagtttaat tttetgtatt eattgattt 2100 ttaacagaag aagaagag

<210> 21

<211> 4368

<212> DNA

<213> Plant

### <400> 21

gaatteetta teeggattte tggtaegeag actgtaatat ggagteatet teteetegat 60 tcgggattaa aattaggtga cttgggacac cctaaatctc ccaagtggcg actctgaaat 120 aaataaacaa atcccgtttc gattgtcctt aaattggaaa aaactccctt gtaccctccc 180 gggtacggaa aaaggaggtg tacagcaatg acccaaaact tttattgcta tacattttga 240 ggaatcaact tgatcaaaat ttatgggtga aattcaatgt ggtatgattt atattaggtc 300 ggactttagc agatgtggtc acttcaattt gcggcaaaaa taatgtacag ggataataat 360 aaaaagtact agaaatttga gtcataaagc tttttcaatt ttacaaaaga tattaagata 420 cttattaaat caaatgtact ttattaatgt aatagcatga aaaaacagcc tcatccgcct 480 gtcctcaccc cacaaaaagg agatagagaa aggaaactaa tcttatttaa ttttccacat 540 tttaatcatc cgtataagaa agaagctaat taactgactt acaaactgaa tagatagcac 660 aatagcacto toaattacaa aaatocaaag oogagggtoa ttootttoat caagaaatta 720 gatagggaat ggaaaatata atttaattat ctgaatcttt ataatttatc cttccatata 780 agaaaaagga aacaaattaa ctgaagagca tatagcctcg catagattta ccttctccat 840 atgggtgggg aaaccgacaa accgcaccaa ttcgataatt cgagtcaaac tgaggaaaaa 900 aaaattcgac tatggtttgg tttgatttgg tttattgttg ggataaaaaa tcgatcataa 960 ttggtttggt ttggttttaa ctaaagaaag tcaaaccgaa accaaacaaa cccgacatta 1020 catatataaa ttttttagat atatttaata tataaatata cttgttgtga tgtaatttat 1080 aaatatttct taaaaatatt cataatttta tcttttaaga tattatttcg tacttagaac 1140 ttttgaatgt ttcttactcc tctttaattt gttagaaatt aatttgaagg agtttgaatt 1200 tgctccgcc ccgctccgtc ccgttgccat ccctgactca ataggataac agcaatctcg 1260 attgcaaaaa tccaaaccca aggaaccttc ccaacattac ataagctaca aagtagagta 1320 gtttattaaa taactaccaa tatatcctca aattctcgcg attatttcat acctaacacg 1380 cttaccttat cttctcgtaa tgacgctaca ttagttggtg atataaaata ccgaatttgc 1440 cacgcggcaa tcctccgctg tctatccacg gcccgagaga atctcttagc cccccaaaga 1500 tgaaaattaa cttctagaat tttatttct ggttattacc atgaaaataa ttaaataaaa 1560 aaaaagagaa aagtaaagat atttaattgg gctaaaactg gggtccacgg cccagccacg 1620

catttccctc	ctatataaag	cgtcgtcacc	tctcatgcaa	atctcgctca	ctacacagtt	1680
gttagtttca	cgttctcttc	tcaattccca	taacagaaac	ccttccgtta	ggtttccgtc	1740
ctattttcc	tcatcttctc	cgtttcctct	tctgaaatca	atatctgtat	ggtgttttc	1800
ttgttcgaat	tttagatttg	ttttgtcttt	aatacctata	accttaaatt	ctctgtttaa	1860
accaaaaact	tagcttcttc	tgaagtcagg	gtggggattt	ttggatcgtg	taagagtgtg	1920
ttagagggtg	attatctttt	gattcagttc	cttttttgct	tcttttgagg	gggtagccgg	1980
ggcctcggcc	tcggcgggtt	ttaatagccc	ccatctatta	caactattgg	gcaaaaacat	2040
cattaaatct	gtacaaaaca	aacccttaat	ttagtttaat	tttctgtatt	cattgatttt	2100
ttaacagaag	aagaagagat	gccggcccta	ggttgttgtg	tagatgctgc	tgttgtttcc	2160
cctcctcta	gctatgcctt	ctctcgggat	agctctcttc	ccgcgccgga	gttctttgcc	2220
tccggcgtac	ctccgacaaa	ttctgccgct	gcttccattg	ggtctccgga	tttgtcgtct	2280
gctttatacg	gggtcgatgg	gtggggagct	ccttatttct	ctgttaactc	taatggagat	2340
atctccgtcc	gaccacacgg	tacggacact	ctccctcacc	aggaaattga	ccttctcaag	2400
gtcgtgaaaa	aggcctccga	cccgaaaaat	tcaggtgggc	ttgggcttca	gctgcctctt	2460
gttgttcgct	tccctgatgt	gttgaaaaac	cggttggaat	ctctgcaatc	ggcttttgat	2520
ctcgcggttc	attcccaggg	ctatggggcc	cactaccaag	gtgtttatcc	cgtgaaatgc	2580
					ccggttcggg	
					caagggcagt	
gctgagggcc	ttctcgtttg	caatggtttc	aaggacgctg	agtacatttc	gcttgctttg	2760
					ggagcttgac	
					tcgggctaag	
					taagtttggg	
					aatgctggat	
					gttgctagct	
					agcgggtatg	
					atcatgcgat	
					tcaggcggtt	
					aagtggcagg	
					tagtcactca	
					gctcaacgaa	
					gtatgagaca	
					agaagggtcc	
					aaaggctatg	
					ttcaattcct	
					agatgaaaag	
					ggttgataag	
					cgatggtggt	
					aggactccac	
aacctgtttg	gtggaccaag	tgtcgtgcgc	gtggtgcaga	gcgatagcgc	tcacagcttt	3960
gccatgactc	gctccgtccc	tggcccgtct	tgcgctgatg	tgctccgagc	gatgcagcac	4020
gagcccgagc	tcatgttcga	gactctcaag	caccgtgcgg	aggaattett	ggaacaagaa	4080
					cttccataac	
					taacaatggt	
					ggaggatgag	
					agtttgtcgt	
cgaggttgtc	tgtttttgaa	taataccctt	agttggtgat	gtttttct		4368

<211> 721

<212> PRT

<213> Plant

<400> 22

Met Pro Ala Leu Gly Cys Cys Val Asp Ala Ala Val Val Ser Pro Pro 1 5 10 15

Leu Ser Tyr Ala Phe Ser Arg Asp Ser Ser Leu Pro Ala Pro Glu Phe 20 25 30

Phe Ala Ser Gly Val Pro Pro Thr Asn Ser Ala Ala Ala Ser Ile Gly 35 40 45

Ser Pro Asp Leu Ser Ser Ala Leu Tyr Gly Val Asp Gly Trp Gly Ala 50 55 60

Pro Tyr Phe Ser Val Asn Ser Asn Gly Asp Ile Ser Val Arg Pro His 65 70 75 80

Gly Thr Asp Thr Leu Pro His Gln Glu Ile Asp Leu Leu Lys Val Val 85 90 95

Lys Lys Ala Ser Asp Pro Lys Asn Ser Gly Gly Leu Gly Leu Gln Leu 100 105 110

Pro Leu Val Val Arg Phe Pro Asp Val Leu Lys Asn Arg Leu Glu Ser 115 120 125

Leu Gln Ser Ala Phe Asp Leu Ala Val His Ser Gln Gly Tyr Gly Ala 130 135 140

His Tyr Gln Gly Val Tyr Pro Val Lys Cys Asn Gln Asp Arg Phe Val 145 150 155 160

Val Glu Asp Ile Val Lys Phe Gly Ser Pro Phe Arg Phe Gly Leu Glu 165 170 175

Ala Gly Ser Lys Pro Glu Leu Leu Leu Ala Met Ser Cys Leu Cys Lys 180 185 190

Gly Ser Ala Glu Gly Leu Leu Val Cys Asn Gly Phe Lys Asp Ala Glu 195 200 205

Tyr Ile Ser Leu Ala Leu Val Ala Arg Lys Leu Met Leu Asn Thr Val 210 215 220

Ile Val Leu Glu Glu Glu Glu Leu Asp Leu Val Ile Asp Ile Ser

225	230	235	240

His Lys Met Ala Val Arg Pro Val Ile Gly Leu Arg Ala Lys Leu Arg 245 250 255

Thr Lys His Ser Gly His Phe Gly Ser Thr Ser Gly Glu Lys Gly Lys
260 265 270

Phe Gly Leu Thr Thr Gln Ile Val Arg Val Val Lys Lys Leu Glu 275 280 285

Glu Ser Gly Met Leu Asp Cys Leu Gln Leu Leu His Phe His Ile Gly 290 295 300

Ser Gln Ile Pro Ser Thr Gly Leu Leu Ala Asp Gly Val Gly Glu Ala 305 310 315 320

Ala Gln Ile Tyr Cys Glu Leu Val Arg Leu Gly Ala Gly Met Lys Phe 325 330 335

Ile Asp Ile Gly Gly Gly Leu Gly Ile Asp Tyr Asp Gly Thr Lys Ser 340 345 350

Cys Asp Ser Asp Val Ser Val Gly Tyr Gly Ile Gln Glu Tyr Ala Ser 355 360 365

Ala Val Val Gln Ala Val Gln Tyr Val Cys Asp Arg Lys Gly Val Lys 370 375 380

His Pro Val Ile Cys Ser Glu Ser Gly Arg Ala Ile Val Ser His His 385 390 395 400

Ser Ile Leu Ile Phe Glu Ala Val Ser Ala Ser Ser His Ser Cys Ser 405 410 415

Ser Ser His Leu Ser Ser Gly Gly Leu Gln Ser Met Ala Glu Thr Leu 420 425 430

Asn Glu Asp Ala Leu Ala Asp Tyr Arg Asn Leu Ser Ala Ala Ala Val 435 440 445

Arg Gly Glu Tyr Glu Thr Cys Val Leu Tyr Ser Asp Gln Leu Lys Gln 450 455 460

Arg Cys Val Asp Gln Phe Lys Glu Gly Ser Leu Gly Ile Glu His Leu 465 470 475 480

Ala Ala Val Asp Ser Ile Cys Asp Phe Val Ser Lys Ala Met Gly Ala

				485					490					495	
Ala	Asp	Pro	Val 500	Arg	Thr	Tyr	His	Val 505	Asn	Leu	Ser	Ile	Phe 510	Thr	Ser
Ile	Pro	Asp 515	Phe	Trp	Ala	Phe	Gly 520	Gln	Leu	Phe	Pro	Ile 525	Val	Pro	Ile
His	Arg 530	Leu	Asp	Glu	Lys	Pro 535	Ala	Val	Arg	Gly	Ile 540	Leu	Ser	Asp	Leu
Thr 545	Cys	Asp	Ser	Asp	Gly 550	Lys	Val	Asp	Lys	Phe 555	Ile	Gly	Gly	Glu	Ser 560
Ser	Leu	Pro	Leu	His 565	Glu	Leu	Gly	Ser	Asn 570	Gly	Asp	Gly	Gly	Gly 575	Tyr
Tyr	Leu	Gly	Met 580	Phe	Leu	Gly	Gly	Ala 585	Tyr	Glu	Glu	Ala	Leu 590	Gly	Gly
Leu	His	Asn 595	Leu	Phe	Gly	Gly	Pro 600	Ser	Val	Val	Arg	Val 605	Val	Gln	Ser
Asp	Ser 610	Ala	His	Ser	Phe	Ala 615	Met	Thr	Arg	Ser	Val 620	Pro	Gly	Pro	Ser
Cys 625	Ala	Asp	Val	Leu	Arg 630	Ala	Met	Gln	His	Glu 635	Pro	Glu	Leu	Met	Phe 640
Glu	Thr	Leu	Lys	His 645	Arg	Ala	Glu	Glu	Phe 650	Leu	Glu	Gln	Glu	Asp 655	Asp
Lys	Gly	Leu	Ala 660	Val	Glu	Ser	Leu	Ala 665	Ser	Ser	Val	Ala	Gln 670	Ser	Phe
His	Asn	Met 675	Pro	Tyr	Leu	Val	Ala 680	Pro	Ser	Ser	Cys	Arg 685	Phe	Thr	Ala
Ala	Thr 690	Asp	Asn	Asn	Gly	Gly 695	Tyr	Asn	Tyr	Tyr	Туг 700	Ser	Asp	Glu	Asn
Ala 705	Ala	Asp	Ser	Ala	Thr 710	Gly	Glu	Asp	Glu	Ile 715	Trp	Ser	Tyr	Cys	Thr 720

Ala

<210> 23 <211> 2695

<212> DNA

<213> Plant

<400> 23

ttcacgttct cttctcaatt cccataaaag aaacccttcc gttaggtttc cgtcctattt 60 tetettette taegetteet ettetgatat caatatetgt atggtgtttt tettgttega 120 attttagatt tgttttgcct ttaatacctg taaccttata attctctgtt taaaccaaaa 180 acttagcttc ttctgaagtc agggtgggga tatttggatc gtgtaagagt gtgttagaag 240 gtgattatct tttgattcag ttcctttttt gcttcttttg agggggtagc cggggcctcg 300 gcctcggcgg gttttaatag cccccatcta ttacaaccat tgggcaaaaa catcattaaa 360 tctgtacaaa gcaaaccctt aatttagttt aattttctgt attctttgat tctttaacag 420 aagaagaaga gatgccggcc ctaggttgtt gcgtagacgc tactgtttcc cctcctctcg 480 gctatgcctt ctctcgggat agctctcttc ccgcgccgga gttctttacc tccggcgtac 540 ctcctacaaa ctccgccgcc ggttccattg ggtctccgga tctgtcctct gctttgtacg 600 gggtcgatgg gtggggaget cettatttet cegttaacte taacggagat atetecgtee 660 gaccacatgg tacggacaca ctcccccacc aggaaattga ccttctcaag qtcqtqaaaa 720 aggcctccga cccgaaaaat tcaggggggc tcgggcttca gctgcctctt gttgttcgct 780 tecetgatgt getaaaaaac eggttggaat etetgeaate ggettttgat etegetgtte 840 atteccaggg etatggggee cactaceaag gtgtttatee egtgaaatge aateaagaea 900 ggttcgtggt ggaagatatt gtcaaattcg ggtcgtcatt ccggttcggg ttggaagctg 960 ggtctaaacc cgagctcctg ttagccatga gctgtctctg caggggcagt gctgagggcc 1020 ttctcgtttg caatggtttc aaggacgctg agtacatttc gcttgctttg gttgcaagaa 1080 agctcatgtt aaacactgta attgttcttg aacaagagga ggagcttgac cttgtgattg 1140 atataagccg taagatggct gttcggcccg taattggact tcgggctaag ctcaqgacca 1200 agcattcagg ccattttgga tccacttctg gagaaaaagg taagtttggg cttacaacga 1260 cccaaattgt tcgtgtagtg aagaagctgg aagaatccgg aatgctggat tgccttcagt 1320 tgctgcattt tcacattgga tctcagatcc cttcaacggc gttgcttgct gatggtgttg 1380 gtgaggctgc tcagatttat tgtgaattaa tccgtcttgg tgcgggtatg aagttcattg 1440 atactggagg tgggctcgga attgattatg atggtactaa atcatgtgat tcagatgtct 1500 ctgttggcta tggcattcaa gaatacgcct ccacagttgt ccaggcggtt caatatgttt 1560 gcgaccgtaa gggcgtgaag cacccagtga tttgcagcga aagtggcagg gcaattgttt 1620 ctcatcactc aattctgatt ttcgaagccg tgtctgcttc tagtcactca tgttcttctt 1680 cacatctgtc ttctggtggc ctccaatcca tggcggagac gctcaatgaa gatgcccttg 1740 ctgattaccg caatttatct gctgctgcag ttcgtggaga gtacgagacg tgtgtacttt 1800 actctgatca gttgaaacag agatgtgtgg atcagtttaa agaagggtcc ttgggtattg 1860 aacatcttgc tgctgttgat agcatctgtg attttgtatc aaaggctatg ggggctgctg 1920 atcctatccg cacttaccat gtgaatctgt caattttcac ttcaattcct gatttttggg 1980 cctttggtca attgtttccg attgttccaa tacaccgttt agatgaaaag cctgcagtaa 2040 ggggaatatt atcggacttg acttgtgaca gtgatgggaa ggttgataag ttcattggtg 2100 gcgaatcaag cttgcagctg catgaattgg gaagtaatgg cgatggtggt gggtattatc 2160 tggggatgtt tttgggtggg gcttatgagg aggcgctcgg aggactccac aacctgtttg 2220 gtggaccaag cgtggtgcgc gtggtgcaga gcgatagcgc tcacagcttc gccatgtctc 2280 gctccgtccc tggcccgtcc tgcgcggacg tgctccgagc gatgcagcac gagcccgagc 2340 tcatgttcga gactctcaag caccgtgcgg aggaattctt ggaacaagaa gaagacaaag 2400 ggctggccat tgcatctttg gccagcagct tagctcagtc cttccataac atgccttacc 2460 ttgtggcgcc tgcatcttgc tgcttcactg cagttactgc taacaacggt ggctataact 2520

actattacag tgatgagaat gcagcagatt ctgctacagg ggaggatgag atttggtcct 2580 attgcactgc ttgaagtgtt gtcgtagcat ctccagtttt agtttgtcgt cgaagttgtc 2640 tgtttttgaa taataccctt agttggtgat gtttttctaa aaaaaaaaa aaaaa 2695

<210> 24

<211> 720

<212> PRT

<213> Plant

<400> 24

Met Pro Ala Leu Gly Cys Cys Val Asp Ala Thr Val Ser Pro Pro Leu

1 5 10 15

Gly Tyr Ala Phe Ser Arg Asp Ser Ser Leu Pro Ala Pro Glu Phe Phe 20 25 30

Thr Ser Gly Val Pro Pro Thr Asn Ser Ala Ala Gly Ser Ile Gly Ser 35 40 45

Pro Asp Leu Ser Ser Ala Leu Tyr Gly Val Asp Gly Trp Gly Ala Pro 50 55 60

Tyr Phe Ser Val Asn Ser Asn Gly Asp Ile Ser Val Arg Pro His Gly 65 70 75 80

Thr Asp Thr Leu Pro His Gln Glu Ile Asp Leu Leu Lys Val Val Lys 85 90 95

Lys Ala Ser Asp Pro Lys Asn Ser Gly Gly Leu Gly Leu Gln Leu Pro 100 105 110

Leu Val Val Arg Phe Pro Asp Val Leu Lys Asn Arg Leu Glu Ser Leu 115 120 125

Gln Ser Ala Phe Asp Leu Ala Val His Ser Gln Gly Tyr Gly Ala His 130 135 140

Tyr Gln Gly Val Tyr Pro Val Lys Cys Asn Gln Asp Arg Phe Val Val 145 150 155 160

Glu Asp Ile Val Lys Phe Gly Ser Ser Phe Arg Phe Gly Leu Glu Ala 165 170 175

Gly Ser Lys Pro Glu Leu Leu Leu Ala Met Ser Cys Leu Cys Arg Gly 180 185 190

Ser Ala Glu Gly Leu Leu Val Cys Asn Gly Phe Lys Asp Ala Glu Tyr 195 200 205

Ile Ser Leu Ala Leu Val Ala Arg Lys Leu Met Leu Asn Thr Val Ile Val Leu Glu Glu Glu Glu Leu Asp Leu Val Ile Asp Ile Ser Arg Lys Met Ala Val Arg Pro Val Ile Gly Leu Arg Ala Lys Leu Arg Thr Lys His Ser Gly His Phe Gly Ser Thr Ser Gly Glu Lys Gly Lys Phe Gly Leu Thr Thr Gln Ile Val Arg Val Val Lys Lys Leu Glu Glu Ser Gly Met Leu Asp Cys Leu Gln Leu Leu His Phe His Ile Gly Ser Gln Ile Pro Ser Thr Ala Leu Leu Ala Asp Gly Val Gly Glu Ala Ala Gln Ile Tyr Cys Glu Leu Ile Arg Leu Gly Ala Gly Met Lys Phe Ile Asp Thr Gly Gly Leu Gly Ile Asp Tyr Asp Gly Thr Lys Ser Cys Asp Ser Asp Val Ser Val Gly Tyr Gly Ile Gln Glu Tyr Ala Ser Thr Val Val Gln Ala Val Gln Tyr Val Cys Asp Arg Lys Gly Val Lys His Pro Val Ile Cys Ser Glu Ser Gly Arg Ala Ile Val Ser His His Ser Ile Leu Ile Phe Glu Ala Val Ser Ala Ser Ser His Ser Cys Ser Ser Ser His Leu Ser Ser Gly Gly Leu Gln Ser Met Ala Glu Thr Leu Asn Glu Asp Ala Leu Ala Asp Tyr Arg Asn Leu Ser Ala Ala Ala Val Arg Gly Glu Tyr Glu Thr Cys Val Leu Tyr Ser Asp Gln Leu Lys Gln Arg 

Cys 465	Val	Asp	Gln	Phe	Lys 470	Glu	Gly	Ser	Leu	Gly 475	Ile	Glu	His	Leu	Ala 480
Ala	Val	Asp	Ser	Ile 485	Cys	Asp	Phe	Val	ser 490	Lys	Ala	Met	Gly	Ala 495	Ala
Asp	Pro	Ile	Arg 500	Thr	Tyr	His	Val	Asn 505	Leu	Ser	Ile	Phe	Thr 510	Ser	Ile
Pro	Asp	Phe 515	Trp	Ala	Phe	Gly	Gln 520	Leu	Phe	Pro	Ile	Val 525	Pro	Ile	His
Arg	Leu 530	Asp	Glu	Lys	Pro	Ala 535	Val	Arg	Gly	Ile	Leu 540	Ser	Asp	Leu	Thr
Cys 545	Asp	Ser	Asp	Gly	Lys 550	Val	Asp	Lys	Phe	Ile 555	Gly	Gly	Glu	Ser	Ser 560
Leu	Gln	Leu	His	Glu 565	Leu	Gly	Ser	Asn	Gly 570	Asp	Gly	Gly	Gly	Туг 575	Tyr
Leu	Gly	Met	Phe 580	Leu	Gly	Gly	Ala	Tyr 585	Glu	Glu	Ala	Leu	Gly 590	Gly	Leu
His	Asn	Leu 595	Phe	Gly	Gly	Pro	Ser 600	Val	Val	Arg	Val	Val 605	Gln	Ser	Asp
Ser	Ala 610	His	Ser	Phe	Ala	Met 615	Ser	Arg	Ser	Val	Pro 620	Gly	Pro	Ser	Cys
Ala 625	Asp	Val	Leu	Arg	Ala 630	Met	Gln	His	Glu	Pro 635	Glu	Leu	Met	Phe	Glu 640
Thr	Leu	Lys	His	Arg 645	Ala	Glu	Glu	Phe	Leu 650	Glu	Gln	Glu	Glu	Asp 655	Lys
Gly	Leu	Ala	Ile 660	Ala	Ser	Leu	Ala	Ser 665	Ser	Leu	Ala	Gln	Ser 670	Phe	His
Asn	Met	Pro 675	Tyr	Leu	Val	Ala	Pro 680	Ala	Ser	Cys	Cys	Phe 685	Thr	Ala	Val
Thr	Ala 690	Asn	Asn	Gly	Gly	Tyr 695	Asn	Tyr	Tyr	Tyr	Ser 700	Asp	Glu	Asn	Ala
Ala 705	Asp	Ser	Ala	Thr	Gly 710	Glu	Asp	Glu	Ile	Trp 715	Ser	Tyr	Cys	Thr	Ala 720

```
<210> 25
<211> 914
<212> DNA
<213> Plant
<400> 25
aagctcgann ttaancctca ntaaagggaa caaaagctgg taccgnggcc cccctcgag 60
gtcgacggta tcgataagct tgattaagct tagtangcac attagcagcg cttgggatga 120
ttttaggcgc ggcctattcc ctttggctat ataatcgtgt ggttctggga attaaaaccc 180
qatttcctcc ataaattctc cgatctaaat ggcanagaat tttcatattt ataccttttc 240
ttgttggagt tgtttggatg ggtgtttacc ccaaagtgtt cctgggactg catgcataca 300
tccqtaaqta acttaaqtqc aacatqqaaa atttcattqa gaggaatcag caaaaaaaaa 360
aagcttaatc gaattcctgc agcccggggg atccactagt tctagagcgg ccgccaccgc 420
ggtggagctc caattcgccc tatagtgagt cgtattacaa ttcactgggc cgtcgtttta 480
caacgtcgtg actgggaaan gcctggcgtt accaacttaa tcgccttgca gcacatcccc 540
ctttcgccag ctgggcgtaa tagcgaanag gccgcacgat cgccttccca acagttgcgc 600
acctgatggn gaatgggacn gecetgtane ngegeangaa negeggengg tgtggtta 660
ccccancgtg acgcnaactt gcaacgccta acgccgcnct tcgcttctcc ttcttctngc 720
aagttcncgg ctttccgtaa gtccaagcgg gggnccttag gttcgattat gttnggaccc 780
ccccnaaact gttanggtnt gtgatttggc accccaaaac gtttcccttg nctggtcntt 840
ttaaangcct ntcaacngaa accaccatcg gnacttgtta aggntncctt gcntgnaaaa 900
                                                                  914
nngtaaatta nntn
<210> 26
<211> 829
<212> DNA
<213> Plant
<400> 26
agcaagctcg atatcgccct cactaaaggg aacaaaaact ggtaccgggc ccccctcga 60
ggtcgacggt atcgataagc ttgattaanc ttttttttt tgaactacac aagggaattt 120
cttctcctna gtaacacatg agaataatta gtgcaataaa ttacaagagg aacattgcag 180
ttggatttaa gaatctgcgc tggggaattt agcctcaata tttgctacaa ccgtacagat 240
ttcactgcat tcatgaacga tagtatccgt gacacatcct tttggatgcc gtcctgtcca 300
catatqccac tactcacatc cactccattg ggtttaagtt gcagaaagag cttcacaaac 360
attctccqqq ttaattcctc ctgccaagag ccacccatgt ttgcttctaa tgcgcggcag 420
cttaaactga acccagttga atcctttgcc actgccacct tttgcactat ccactagaac 480
ccaatcaacc agagaagact cctcatcaga aatatagttc aaaaggctcc tcttcatttg 540
catgaagtac gtatattaca cgtttttccc tgaccaaagc ttaatcgaat tctgcagccc 600
gggggatcng gnattctaga gcggcgccac gcggtggagc tccaatcgcc taaatgancn 660
ataaaatcac tggccgtcgt ttanacgncn ggacgggaaa cctgggtacc aacttaatcg 720
cctgnagcna tccccttcnc agcggngtan acgaaaggcc gncgattgcc tccanattgc 780
                                                                  829
cacnggatgg aanggacncc gtncgganga acngggggnn ggggtaccn
```

International application No.
PCT/US00/12450

IPC(7) :	SSIFICATION OF SUBJECT MATTER A01H 5/00; C07H 21/04; C12N 5/14, 15/29, 15/52, 435/320.1, 414, 419; 536/23.2, 23.6, 24.5; 800/278,	317.3	*				
	o International Patent Classification (IPC) or to both	national classification and IPC					
	DS SEARCHED						
	ocumentation searched (classification system followe	,					
U.S. : 4	435/320.1, 414, 419; 536/23.2, 23.6, 24.5; 800/278, 3	317.3					
Documentat	ion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)							
Please See	Please See Extra Sheet.						
C. DOC	UMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.				
X	HASHIMOTO et al. Intraspecific Varia	ability of the Tandem Repeats	12				
	in Nicotiana Putrescine N-methyltr	ansferase. Plant Molecular					
Y	Biology. 1998, Vol. 37, pages 25-37,	, especially Figure 3.	15,16				
X	HIBI et al. Gene Expression in Tobacc	o Low-Nicotine Mutants. The	12				
	Plant Cell. May 1994, Vol. 6, pages 7						
Y			15,16				
X	IZHAKI et al. A Petunia cDNA End	coding S-Adenosylmethionine	12				
	Synthetase. Plant Physiology. 1995, V	,					
Y	entire article.	, F. B	15,16				
			,				
İ							
X Furth	er documents are listed in the continuation of Box C	. See patent family annex.					
• Spe	cial categories of cited documents:	"T" later document published after the inte					
	nument defining the general state of the art which is not considered be of particular relevance	date and not in conflict with the appli the principle or theory underlying the	invention				
	lier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be consider	ed to involve an inventive step				
cite	nument which may throw doubts on priority claim(s) or which is do establish the publication date of another citation or other	when the document is taken alone  "Y" document of particular relevance; the	ataba at barantina samust be				
•	cial reason (as specified)  cument referring to an oral disclosure, use, exhibition or other	considered to involve an inventive combined with one or more other such being obvious to a person skilled in the	step when the document is documents, such combination				
"P" doc	nument published prior to the international filing date but later than priority date claimed	"&" document member of the same patent					
Date of the	actual completion of the international search	Date of mailing of the international see	rch report				
17 AUGU	ST 2000	0 4 001 200	U				
	nailing address of the ISA/US	Authorized officer	8010				
Box PCT	ner of Patents and Trademarks	AMY NELSON	(XXXXXII)				
	, D.C. 20231 b. (703) 305-3230	Telephone No. (703) 308-0196					

International application No.
PCT/US00/12450

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
X  Y	LAMATTINA et al. RNA Editing of the Transcript Coding for Subunit 4 of NADH Dehydrogenase in Wheat Mitochondria: Uneven Distribution of the Editing Sites Among the Four Exons. Nucleic Acids Research 1991, Vol. 19, No. 12, pages 3275-3282, especially Figure 4.	12  15,16
X  Y	LI et al. Arabidopsis Phosphoribosylanthranilate Isomerase: Molecular Genetic Analysis of Triplicate Tryptophan Pathway Genes. The Plant Cell. April 1995, Vol. 7, pages 447-461, especially Figure 3, page 459.	12,15  16

International application No.
PCT/US00/12450

Bo	x I C	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)							
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:									
1.		Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:							
2.		Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:							
3.		Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).							
Во	x II (	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)							
Thi	is Inter	mational Searching Authority found multiple inventions in this international application, as follows:							
	Ple	ease See Extra Sheet.							
1.		As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.							
2.		As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.							
3.	X 1-	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: -15,18-20							
4.		No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:							
Re	mark (	on Protest  X The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.							

International application No. PCT/US00/12450

#### **B. FIELDS SEARCHED**

Electronic data bases consulted (Name of data base and where practicable terms used):

STN, AGRICOLA, CAPLUS, BIOSIS, EMBASE, USPAT

search terms: putrescine methyltransferase, adenosylmethionine synthetase, omithine decarboxylase, arginine decarboxylase, NADH dehydrogenase, phosphoribosylanthranilate isomerase, DNA, cDNA, gene, nucleic

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-16, drawn to coding DNA, vector, host cell, transgenic plant.

Group II, claim(s) 17, drawn to protein.

Group III, claim(s) 18-20, drawn to transformation method and transgenic plant with promoter DNA.

The inventions listed as Groups I, II, and III do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The coding DNA of Group I, e.g. Claim 12, is disclosed in the prior art publication of Hashimoto et al. (Plant Mol. Biol. 37: 25-37, 1998; see Fig. 3b). Therefore, there is no special technical feature which links the coding DNA of Group I with the protein of Group II.

Furthermore, there is no special technical feature under PCT Rule 13.2 which links the coding DNA of Group I and the transformation method and transgenic plant with the promoter DNA of Group III. Therefore, the inventions of Groups I, II, and III do no relate to a single inventive concept under PCT Rule 13.1.